A Search For New Low-Mass Diphoton Resonances At Atlas And An Investigation Into Using Gaussian Process Regression To Model Non-Resonant Two-Photon Standard Model Backgrounds

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Abstract
The Standard Model of particle physics has been tested over many years with many experiments and has predicted experimental results with remarkable accuracy. In 2012, the last piece of the Standard Model, the Higgs boson, was discovered by the experiments ATLAS and CMS at the Large Hadron Collider (LHC). Although this completes the Standard Model, this by no means completes our picture of the physics that describes the observable universe. Several phenomena and measurements remain unexplained by the Standard Model including gravity, dark matter, the baryon-antibaryon asymmetry of the universe and more. One of the primary goals of the LHC and the ATLAS experiment are to search for extensions and modifications to the Standard Model that could help to explain these phenomena. This thesis presents three areas where I made major contributions. The first is in the identification of prompt electrons in ATLAS using a likelihood method both in the online trigger system and in offline data analysis. Prompt electrons are ubiquitous in the signatures of electroweak physics, one of the cornerstones of the ATLAS physics program. Next I present a search for new physics in low-mass (65-110 GeV) diphoton events. This is a model independent search that is motivated by several extensions to the Standard Model including the two Higgs doublet model where new scalars can appear as lighter versions of the Standard Model Higgs. No evidence for a new narrow resonance is found, so limits ranging from 30 to 101 fb are set on the production cross section of such a resonance, assuming that its branching fraction to two photons is 100 percent. The sensitivity of these results are limited by the systematic uncertainties due to the potential spurious signals introduced by the two-photon non-resonant Standard Model background. My third contribution was an initial investigation of a new method to model this background using Gaussian Process Regression.

Degree Type
Dissertation

Degree Name
Doctor of Philosophy (PhD)

Graduate Group
Physics & Astronomy

First Advisor
I. J. Kroll

Keywords
Diphoton

Subject Categories
Physics

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A SEARCH FOR NEW LOW-MASS DIPhotON RESONANCES AT ATLAS AND AN
INVESTIGATION INTO USING GAUSSIAN PROCESS REGRESSION TO MODEL
NON-RESONANT TWO-PHOTON STANDARD MODEL BACKGROUNDS

Rob Roy MacGregor Fletcher

A DISSERTATION
in
Physics and Astronomy

Presented to the Faculties of The University of Pennsylvania
in Partial Fulfillment of the Requirements for the Degree of Doctor of Philosophy
2018

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A SEARCH FOR NEW LOW-MASS DIPHOTON RESONANCES AT ATLAS AND AN INVESTIGATION INTO USING GAUSSIAN PROCESS REGRESSION TO MODEL NON-RESONANT TWO-PHOTON STANDARD MODEL BACKGROUND

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Preface

This thesis describes some of the work I did as a graduate student but does not attempt to be a complete and detailed discussion of any topics. My hope is that someone will find this useful, especially the last chapter which describes my attempts to refine how background modeling in the low mass diphoton group is done. This work will likely be carried on by new students and this may be a good starting point to see what I have tried in the past, what work I was doing at the end, and what studies were outstanding. This will also be a source of references that describe in much more detail the topics contained in here.

This thesis will be organized as follows: Chapter 1 contains an overview of my time in graduate school, with very brief descriptions of work I performed that are not the focus of later sections. This section is largely historical and is intended to document projects for my own benefit. Chapter 2 gives a short introduction to the Standard Model of particle physics as well as a few shortcomings of the theory. As the main analysis described in this thesis is motivated by extensions to the scalar sector of the Standard Model, emphasis is placed on how additional scalars can help to address gaps in our understanding of the fundamental particles and their interactions. Chapter 3 describes the Large Hadron Collider and the ATLAS experiment. This chapter skips over a tremendous amount of detail, but references give complete descriptions of both the LHC and ATLAS. Each collaborator on the ATLAS experiments earns the right to be an author on physics publications by performing a qualification task. My task was
performed in a so-called e/gamma group, a Combined Physics and Performance group that focuses on the identification of electrons and photons whose experimental signatures are very similar in the ATLAS detector. I worked on the identification of prompt electrons produced in the decay of electroweak processes such as $W \rightarrow e\nu$, $Z \rightarrow e^+e^-$. This work is described in Chapter 4. After qualifying for authorship, I began the work that would make up the bulk of this thesis. Experimental signatures with two photons in the final state provide clean processes with relatively small backgrounds which makes them a good choice to search for new physics. A search for low mass diphoton resonances below the Standard Model Higgs mass is described in Chapter 5. The analysis on diphoton events uses a method known as spurious signal to test for functional forms to describe the background distribution. This method requires large quantities of simulated data which are prohibitively expensive to generate. Chapter 6 describes some initial investigation with the goal to replace this method with a more advanced technique based on Gaussian Process Regression. This method was ultimately not used in this round of analysis, however I think is a very promising way forward for not only the diphoton group, but many other groups with similar regression tasks.

"Sometimes science is more art than science, Morty. A lot of people don’t get that."

– Rick Sanchez

Rob Roy MacGregor Fletcher

University of Pennsylvania, July 2018
CHAPTER 1

Introduction

The discovery of new particle consistent with the Higgs boson was announced on July 4th, 2012. During the announcement, I was driving a truck through western Pennsylvania on my way to start graduate school at Penn. This was an exciting time in particle physics, and Penn had played a big part. The summer before classes began for me, I worked with Doug Schaefer, one of the Penn graduate students at that time, on the WW decay channel of the Higgs boson. Although I did not work on this for too long, it got me excited to finish classes and dig into more analysis.

This chapter will briefly describe some of the work that I did on ATLAS that does not fit well into other sections or does not merit its own section. This work is primarily being documented for my own benefit.

1.1 Electron ID

My first real work on ATLAS was in the Egamma group which is a combined physics and performance group that focuses on the identification of prompt electrons and photons. My focus was the identification of electrons. This was to be my service work on ATLAS. When I joined the group, the electron ID had been reworked by a few Penn students and had been
changed from a cuts-based system, to a naive Bayes method called electron likelihood. At this time the likelihood approach was only used in the offline analysis; the cuts-based analysis had been used in the trigger. There was interest in moving the trigger over to an electron likelihood as well. This would remove some inefficiencies that arose from using 2 different IDs. Some electrons that passed the cuts ID would not pass the likelihood and vice versa. Work began to modify the likelihood to work in the trigger. For me this culminated in 2 sleepless weeks with another Penn student Joey Reichert and I trying to finalize new versions of the likelihood before the deadline to get them into the trigger.

1.2 Large Extra Dimension Models

After my service work was done, I transitioned into the exotic diphoton group. This group focuses on searches for exotic particles whose final states include two photons. In the beginning I was interested in graviton models that could be measured in the diphoton channel. The analysis I had started on was a search for Arkani-Hamed, Dimopoulos and Dvali (ADD) type gravitons \cite{1} and Randall-Sundrum type gravitons \cite{2}. These models are also sometimes known as Large Extra Dimension (LED) models as they propose an additional spatial dimension, $x^5$. If we consider a periodic extra dimension,

$$x^5 \sim x^5 + 2\pi R$$

where $R$ is the radius of extra dimension, the momentum of a particle propagating in this dimension would be

$$p^5 \sim \frac{n}{R}, \quad n \in \mathbb{Z}.$$ 

Then we can write the equation motion of such a particle as,

$$\delta_{\mu} \delta^{\nu} \phi^n(x^\mu) = \frac{n^2}{R^2} \phi^n(x^\mu)$$
1. Introduction

with mass,

\[ m^2 = \frac{n^2}{R^2}. \quad (1.4) \]

Theories of this type with an extra dimension are known as Kaluza-Klein theories after Theodore Kaluza and Oscar Klein. These theories have been tested down to extremely small scales with current limits at \( R \lesssim 10^{-21} \) meters. Extra dimensions of this size are pretty much hopeless to find. Arkani-Hamed, Dimopoulos and Dvali proposed that maybe this extra dimension is only accessible to gravity and, this causes the size of the extra dimension to only be bounded by tests of gravity, which means \( R \lesssim 1 \) mm. One very interesting feature of theories like this is that since only gravity can propagate through this extra dimension, it can have the effect of “diluting” the effect of gravity in the 3 other space dimensions. This can possibly explain why there is such a large difference between the strength of the weak force and gravity. This difference is known as the hierarchy problem.

What does gravity look like in more than the 3 dimensions we are used to? The gravitational potential we all learn in our beginning physics classes can be written,

\[ V(r) = G\frac{m_1m_2}{r} \quad (1.5) \]

where \( G \) is the gravitational constant and \( r \) is the distance between masses \( m_1 \) and \( m_2 \). Using the relationship \( M_{pl} \sim \sqrt{\frac{1}{G}} \) (where I’m leaving out factors of \( hc \) and \( \pi \) for simplicity) where \( M_{pl} \) is the Planck mass, we can write this as

\[ V(r) \sim \frac{1}{M_{pl}^2} \frac{m_1m_2}{r} \quad (1.6) \]

Now, if we add in \( n \) extra dimensions, we can use Gauss’s law, considering \( r \ll R \), to write the gravitational potential as,

\[ V(r) \sim \frac{1}{M_{D}^{2+n}} \frac{m_1m_2}{r^{1+n}} \quad (1.7) \]
1. Introduction

We can see that if \( n > 0 \) for relatively small scales we should measure the potential deviating from the familiar \( 1/r \) law. What about at scales where gravity has been well measured \( (r \gg R) \)? In this case the potential can be written as,

\[
V(r) \sim \frac{1}{M^{2+n}_P R^n} \frac{m_1 m_2}{r} \tag{1.8}
\]

Comparing equations 1.6 and 1.8 we can see that this relates the 3-dimensional Planck mass that we measure, to the n-dimensional one where gravity is allowed to propagate in more dimensions.

\[
M^2_\text{pl} \sim M^{2+n}_P R^n \tag{1.9}
\]

Using this relationship we can begin to try to explain the hierarchy problem by setting the n-dimensional Planck mass to the weak scale \( \sim 1 \text{ TeV} \). This is saying that there really is no difference in the forces when considering n-dimensions, and we only see an apparent difference because we can only observe the 3-dimensional version of the Planck mass. What does this mean for the radius of the extra dimensions \( R \)? For \( n = 1 \) we get that \( R \sim 10^{13} \text{ cm} \). This is on the order of the size of the solar system! This definitely does not agree with experiment. For \( n = 2 \) we get that \( R \sim 1 \text{ mm} \) which we said was around the smallest scale tests of gravity have been performed. So tests of gravity do not rule out extra dimension models with a number of extra dimensions \( n = 2 \) or greater.

The way that just adding extra spatial dimensions could very simply address the hierarchy problem was very interesting to me. I joined the search for large extra dimensions using data from the early part of Run2 at ATLAS. Results published on 2015 data showed a 3.4\( \sigma \) deviation from Standard Model backgrounds at 750 GeV, with CMS (the competing experiment to ATLAS) also showing a 3.9\( \sigma \) deviation [5, 6]. Although the threshold for discovery used in high energy physics is 5\( \sigma \), both experiments seeing an excess at \( \sim 750 \text{ GeV} \) was enough to spark intense interest from the theoretical community. Around 200 papers were published
within a month of the results being announced. During this time I participated in cross checks of the results [7].

1.3 HGam Data Validation

Early in my time with the exotics diphoton group, it was decided that this group should be merged with the Higgs to gamma gamma group (HGam) as their analysis was very similar and the background samples were largely the same. My first large project in the HGam group was to work on a framework that could perform validation on smaller datasets derived from the larger raw datasets collected by ATLAS. The validations primary goal was to ensure that no data was missing or incorrectly calculated in the derived data. Another Penn student, Tony Thompson, had begun this project and had produced very rough skeleton of a framework when he left the experiment. My main contribution to this was to write a script that automatically checked whether the proper data sets were created and if the information they contained matched the raw data. This was done in python and the outputs were written to JSON. The outputs were then used to build a dynamic website that contained easy to read tables with all of the relevant information so that it could be used as a reference for anyone in the group. Figure 1.1 shows a screen shot of the website. When the script completed running over the datasets it would also email a report summarizing any issues it found to the people responsible for creating the datasets. Although this project was not directly related to Higgs physics, it was very fun and one of the first times I had built a complete website that functioned dynamically based on underlying data.
1.4 High-Mass Diphoton

The high-mass diphoton search in the HGam group included searches for new scalar particles, non-resonant and resonant spin-2 signals in the mass range 200 GeV - 2.7 TeV \[8, 9\]. In order for the results to be reported in a way that is useful to theorists and for comparison to other experiments, they must be corrected to account for various detector and selection effects. For example, if we measure the cross-section of a particle by counting how many times we observe that particle and comparing that with the integrated luminosity collected, this will give incorrect results as some of the particles produced may follow a path that takes it through gaps in the detector, or regions of the detector that are used for cabling or support structure. Or maybe that particle had some kinematic configuration that meant it was rejected by our selection, or by our trigger due to technical or computing limitations. If a theorist wanted to compare our measurement to his or her model, they would need to know the actual...
cross-section, not just the part of it we can see. In order to correct these values we calculate acceptance and correction factors by using Monte Carlo (MC) simulation. To do this, we take generated MC and find the ratio of particles that pass our analysis selection, or that pass through the active volume of the detector to the total number generated. These ratios are shown in equation 1.10

\[ C_X = \frac{N_{\text{selection}}}{N_{\text{fiducial}}} \quad A_X = \frac{N_{\text{fiducial}}}{N_{\text{total}}} \] (1.10)

Here \(N_{\text{selection}}\) is the number of particles passing the applied selection cuts, \(N_{\text{total}}\) is the total number of particles generated in Monte Carlo, and \(N_{\text{fiducial}}\) is the number of particles generated in Monte Carlo that pass through the fiducial volume of the detector. The definition of the fiducial volume of the detector is something that can take a lot of work on the part of analysts to accurately describe. For the sake of simplicity this can be thought of as a region of the detector where its response is well understood and where all components are functioning properly. In the high-mass diphoton search paper released in 2017 [8], I was responsible for calculating these so-called AxCx factors and their associated uncertainties. The factors and the fits used to model them are shown in figure 1.2 To calculate the uncertainties on these factors several modified samples are produced. These samples have some quantity modified away from its nominal value up \(1\sigma\) and down \(1\sigma\). The AxCx factors are recalculated for each of these and their difference is fit with a function. This function is then used to model the uncertainty on the factors due to imperfect knowledge of the quantity being varied. Figure 1.3
1. Introduction

Figure 1.2: The acceptance factor (a) and the correction factor (b) for several resonance widths and at several masses of a scalar particle. The first part of the label Xtoyy refers to MC produced from a new particle X decaying to two photons. The last part of the labels are used to designate the decay width of the particles used in the MC samples, NW being the nominal narrow width sample.

Figure 1.3: The uncertainty and fit function used for the photon ID efficiency. The closed circles represent variation of the nominal value up by 1σ. The open circles represent the variations down by 1σ.
Chapter 2

The Standard Model

The Standard Model of particle physics is the name given to theory used to describe matter and how it interacts. It has taken form over the course of several decades and began with separate theories. The first steps toward a unified picture of the fundamental particles and forces came in 1961 when Sheldon Glashow unified the weak and electromagnetic forces [10]. This led to the current description of the Standard Model that began to solidify in the 1970s with the experimental confirmation of quarks in deep inelastic scattering experiments conducted at the Stanford Linear Accelerator [11].

2.1 The Standard Model

The Standard Model as it is now described is a type of quantum field theory known as a gauge theory whose local transformations obey the symmetry groups,

\[ SU(3)_C \times SU(2)_L \times U(1)_Y \]  

(2.1)

when arranged in particular multiplet representations. Interactions between all of the matter particles are described by gauge bosons which appear as fields that must be added into the field content of the model in order to ensure local invariance under these transformations. This is done by the modification of the derivative to the covariant derivative which depends
2. The Standard Model

on these new gauge fields. The generators of the respective symmetry groups provide the description of the gauge fields responsible for the fundamental forces. The 8 generators of $SU(3)$ are the 8 gluon fields $G_\mu$ of the strong interaction, with the 3 generators of the $SU(2)$ group combined with the single generator of the $U(1)$ group giving the $W_\mu$ and $B_\mu$ fields responsible for the electroweak interactions.

The symmetry groups as written require that all particle content in the Standard Model be massless which is contrary to experiment where the fermions and weak bosons are observed to be massive. The solution to this problem was proposed through the Higgs mechanism developed by three groups: Peter Higgs [12, 13], Robert Brout and Francois Englert [14], and Gerald Guralnik, Carl R. Hagen, and Tom Kibble [15]. This mechanism breaks the electroweak symmetry through the addition of a scalar field with non-trivial vacuum.

$$SU(2)_L \times U(1)_Y \rightarrow U(1)_{EM}$$  \hspace{1cm} (2.2)

After symmetry breaking the scalar field acquires a vacuum expectation value in the low-energy regime and the interaction with the fermionic fields and the gauge fields gives them mass. The $W_\mu$ and $B_\mu$ fields mix to give the physical $W^\pm$ and $Z$ massive gauge bosons which are the mediators of the weak force. The $U(1)_{EM}$ symmetry is responsible for an orthogonal mixing to give the massless gauge field $A_\mu$ of the photon responsible for mediating the electromagnetic force.

The last piece of the standard model was added in the 1970s which described the interactions of the strong interaction in the theory of quantum chromodynamics (QCD). Quarks experience a phenomena known as confinement due to the fact that their interaction becomes stronger and larger distances. This posed a problem for experiment as it predicted that quarks could never be detected on their own but must exist in bound states with other quarks. Politzer, Wilczek and Gross [16, 17, 18] in 1973 showed the asymptotic freedom of
The combination of the electroweak theory and quantum chromodynamics are what became known as the Standard Model of Particle Physics (SM). The field content of the SM is illustrated in figure 2.1.

2.2 Beyond the Standard Model

The predictions made by the Standard Model have agreed remarkably well with experiment. However, we know that the SM cannot be the only component in our description of physics. One of the most glaring shortcomings of the SM is the complete lack of any description of
2. The Standard Model

gavity. There have been many attempts at a quantum theory of gravity and unifications with
the 3 other forces in the SM, but as of yet none have been completely successful. There are
several other places where the SM fails to give any specific reason for some effect we know
exists, or provides no prediction on an experimentally observed quantity. Endeavors to expand
or modify the Standard Model are collectively known as Beyond the Standard Model.

2.2.1 Baryon Asymmetry

In all of the observable universe we see almost entirely matter. We do not seem to see hardly
any antimatter. In the early universe these two should have been created in equal parts and
experiment seems to tell us that they annihilate in equal parts. Something must have caused
the universe to prefer one over the other. In 1967 Sakharov [20] proposed three conditions
that must be met in order to have an asymmetry in baryonic matter in the universe. These
are:

1. A baryon number violating process

2. Charge \((C)\) and Charge-Parity \((CP)\) violation

3. Interactions outside of thermal equilibrium

Baryon number \((B)\) is obviously a requirement as if we assume initially \(B = 0\) and in the
later we observe \(B\) to be a large number, then this number cannot be conserved. There is no
known process in the Standard Model that violates baryon number. \(C\) and \(CP\) violation both
exist in the weak interactions but the amount of violation is not large enough to account for
the remnants of the early universe that we observe today. Several extensions to the Standard
Model address these in different ways. The two Higgs doublet model adds a second scalar
doublet to the SM and allows for more places where \(CP\) violation can occur. Other models
address the asymmetry in a more direct way, for example Darkogenesis\textsuperscript{[21]} where the baryon asymmetry is generated through interaction with a "dark" sector of matter that can in effect transfer baryon number to the visible sector.

### 2.2.2 Dark Matter

Dark matter is another area where the Standard Model remains silent. We know from observations of galactic rotation that the luminous matter cannot account for all of the mass in a galaxy\textsuperscript{[22]}. It is also observed that large regions of space bend light in an effect called gravitational lensing\textsuperscript{[23]}. The standard model does not contain any particle that could exhibit all of the properties of dark matter, the most important of which is that it does not interact with photons (hence the name 'dark').

![Figure 2.2: Rotation curve of the spiral galaxy M33. The yellow and blue points are the measured curve, while the dotted line is the one predicted from luminous matter only\textsuperscript{[24]}](image)

Extensions to the Standard Model need to add a particle that does not couple to photons,
2. The Standard Model

has a very small cross section to any other SM particle, has mass and is stable. Some extensions to the scalar sector of the SM contain what are called inert fields that do not interact with the rest of the SM. These massive bosons could then be a candidate for dark matter. Supersymmetric models also contain particles that can fill this role.

2.2.3 Supersymmetry

There are several search programs looking for new particles beyond the SM. The most popular of these programs is the search for Supersymmetry (SUSY), a type of space-time symmetry that supposes that all fermions have bosonic super partners, and all bosons have fermionic super partners [25]. This approximately doubles the particle content in the SM. In addition, at least one more scalar doublet field must be added. One of Supersymmetry’s biggest draws is that it solves the Hierarchy Problem where divergent terms in the SM cancel against terms in SUSY. This also solves the problem of the unification of forces. Figure 2.3 shows the running couplings of the 3 forces in the SM, and with SUSY. As can be seen the SM couplings never completely unify as all three forces don’t converge at a single point. However when including SUSY, the running couplings are modified and the forces do all unify at a point. Supersymmetry can also provide a candidate for dark matter as the couplings of SUSY particles to the SM are tightly constrained and so can have relatively stable light particles [26]. These particles could easily have couplings to the photon and other SM particles smaller than observed coupling limits on dark matter.
Figure 2.3: The running couplings of the forces in the SM shown with the dotted lines. With the addition of Supersymmetry, the forces have a point of unification at high energy scales, shown in red and blue [25].
The LHC and ATLAS Detector

The Large Hadron Collider (LHC) is one of the largest, most complicated machines ever built by man. It has fascinated me for years and is one of the reasons that I chose to study high energy physics. The detectors built to take advantage of the collisions produced in the LHC are no less impressive and represent the cumulative work of thousands of physicists and engineers. This chapter will briefly introduce the LHC and the ATLAS detector. This is by no means a complete description. For more detail on the LHC see references [27, 28] and for ATLAS see references [29, 30, 31, 32].

3.1 The Large Hadron Collider

The LHC is a particle collider consisting of about 10,000 superconducting magnets. It accelerates protons around the 27 kilometer ring and collides them with a center of mass energy of $\sqrt{s} = 13$ TeV. The facility is built at European Organization for Nuclear Research (CERN) about 100 meters underground across the French-Swiss border. The first collisions were recorded in 2010 with a center of mass energy of $\sqrt{s} = 7$ TeV. Protons collide at four interaction points along the beamline where the four experiments sit. These are ATLAS and CMS, which are general purpose detectors, ALICE and LHCb which are detectors designed
for heavy ion collisions and b-quark physics respectively. The LHC is the final stage of a series of accelerators that raise the beam energy until it can be directed into the main beamline. The CERN accelerator complex is shown in figure 3.2

Protons in the LHC travel around the ring in bunches of $1 \times 10^{11}$ protons per bunch. They circulate at a rate of 50 Mhz (25ns bunch spacing) and each time bunches cross there are approximately 30 interactions under the current run conditions. A quantity called instantaneous luminosity is used to measure the rate that data is taken. Figure 3.3 shows the instantaneous luminosity per day for data taking in 2017. The integrated luminosity is how we measure the amount of data taken and is quoted in units of inverse femtobarns ($fb^{-1}$). These units make it easy to estimate the number of events we can expect for a given process with cross-section $\sigma$ (measured in femtobarns). In the data taking period from 2015 to 2017 $\sim 80 fb^{-1}$ were recorded with the ATLAS detector. Figure 3.4 shows the luminosity recorded by ATLAS per month for the years 2011 through 2018. Although such high luminosities come...
3. The LHC and ATLAS Detector

Figure 3.2: A schematic drawing of the CERN accelerator complex responsible for feeding protons into the main LHC beamline. (Image: CERN)

with challenges due to the rate of interactions, observation of extremely rare processes require that we run at these high rates.

3.2 The ATLAS Detector

The LHC houses two general purpose detectors; ATLAS and CMS. These detectors have been designed to accommodate a very broad physics program and serve as cross checks of each other. In this section I will give a brief introduction to the design of the ATLAS detector with a focus on the detector subsystems that are relevant to electron identification.
3. The LHC and ATLAS Detector

Figure 3.3: Peak luminosity by day for data taking in 2017 [33].

Figure 3.4: The luminosity per month for years 2011-2018 recorded by ATLAS [34].
3. The LHC and ATLAS Detector

3.2.1 Overview

The ATLAS detector is situated at one of the interaction points along the LHC that is on the CERN campus. It is located approximately 100m underground, is 25m high, 44m long and weighs around 7000 tons. Proton interactions take place at the center of the detector and are recorded as the particles created in the interaction escape outward. Figure 3.5: The ATLAS detector is composed of several subsystems that are designed to make different kinds of measurements. These can generally be split into 3 systems; trackers (collectively known as the inner detector), calorimeters, and muon spectrometer. These systems are arranged in a concentric pattern around the interaction point, and is made up of a barrel shaped section that is approximately uniform in the plane orthogonal to the beamline, and end cap sections that are circular and located on either end of the barrel.

Figure 3.5: A cutaway view of the ATLAS detector. (Image: CERN)
3.2.2 Inner Detector

The inner detector is responsible for tracking particle paths as the pass through the detector. The inner detector is composed of 3 separate subsystems; the pixel detector, silicon strip detector (SCT), and the transition radiation tracker (TRT). The arrangement of these subsystems is shown in figures 3.6 and 3.7.

![Figure 3.6: A cross section of the inner detector showing all layers of the subdetector. (Image: CERN)](image)

The pixel detector consists of 80 million channels in 3 layers. Just outside of this is the SCT microstrip detector with 4 layers. These two detectors both use silicon based tracking where charged particles passing through the layers causes current to flow and a corresponding point in space is recorded in data. The third layer works a bit differently to the previous two. The transition radiation tracker consists of straw tubes filled with xenon (or argon in
Figure 3.7: A cutaway of the inner detector showing the arrangement of all subsystems in relation to each-other. (Image: CERN)

some sections) and a wire suspended in the middle. When charged particles pass through the straw tubes the gas inside is ionized and the ions drift to the walls of the straw tube where the total charge collected is recorded. This can serve a tracking function as there are 50,000 straws in the barrel section and the path of the particle can be reconstructed by finding the line of straws that have signal in them. The reason for this design comes from the transition radiation (TR) emitted by particles interacting with detector. Lighter charged particles will emit more transition radiation than heavier ones. This allows the use of the TR to help distinguish electrons from other particles and is a valuable input for electron identification. These three trackers are housed inside of a solenoid magnet with 2 Tesla field. This field causes charged particle tracks to bend and by calculating the radius, the momentum and charge of the particles can be determined.
3.2.3 Electromagnetic Calorimeter

Calorimeters are used to measure the energy deposited in them as particles pass through, or stop in them. The first of these is the Liquid Argon Calorimeter (LAr) also known as the electromagnetic calorimeter (EMcal) [35]. It uses liquid argon as a scintillator and accordion shaped electrodes arranged in 3 layers. The layers are segmented differently to measure the shape of electromagnetic showers in the material. The first layer is very finely segmented in the \( \eta \) direction, the second and deepest layer is segmented evenly in the \( \eta \) and \( \phi \) direction. The third layer is the most shallow and has larger segmentation than the other two. A slice of the EMcal is shown in figure 3.8. The total depth of the EMcal is \( \sim 22 \) radiation lengths so most showers should be entirely contained within this subsystem. This calorimeter is the primary system used in electron and photon identification.

3.2.4 Hadronic Calorimeter

Located just outside of the EMcal is the Hadronic Calorimeter, also called the tile calorimeter (tileCal). Each of 64 modules in the barrel are composed of alternating layers of iron plates and scintillating tiles. As the name suggests, this calorimeter is designed to measure the energy of hadronic particles. The tiles sample the hadronic shower and based scintillating measured the total energy in the shower can be calculated.

3.2.5 Muon Spectrometer

Along with neutrinos, muons are one of the only particles that will normally completely escape from the detector. While other particles are completely stopped in the calorimeters, muons will pass through the outer layers. The muon spectrometer consists of chambers that measure the momentum of muons. The system uses a series of 4 Tesla toroid magnets.
3. THE LHC AND ATLAS DETECTOR

Figure 3.8: A small slice of the ATLAS Electromagnetic Calorimeter system showing how each of the 3 layers is segmented [36].
Figure 3.9: A cutaway view of the calorimeter system in the ATLAS detector. The EM calorimeter and Hadronic calorimeter are shown \cite{37}.
3. The LHC and ATLAS Detector

Figure 3.10: One barrel module of the hadronic calorimeter showing the arrangement of iron and scintillating tiles [38].

Figure 3.11: A cutaway view of the ATLAS muon spectrometer [39].
Electrons are an important part of many ATLAS analyses. Whether directly looking for decays involving electrons or identifying them to veto events, electron ID finds its way into use in a large number of groups. The primary instruments used in ID are the pixel and silicon-strip tracking systems, transition radiation tracker, EM calorimeter and to a lesser extent the hadronic calorimeter. A number of quantities (variables) are measured in each of these subsystems are used to differentiate electrons and photons from other particles. In the past a cuts-based ID was used where each variable is subdivided into an accept and reject region. If an electron candidate lies in the accept region, this process is repeated for the next variable. This continues until all of the variable distributions have been cut on. The candidates that pass all of these cuts are then considered to have passed some level of electron identification. The line dividing the accept and reject regions can be adjusted to control the efficiency of identification for what we would call an operating point. Although this method is simple and provided a fairly small rate of false positive, with the increasing demands placed on the detector due to higher luminosity, a better method was sought. The Electron Likelihood was selected to replace the cuts based method [40][41].

My work on ATLAS began in the Egamma group with the Electron Likelihood. While working on electron ID, I contributed almost all areas of the Likelihood ID, but a majority
of my work involved preparing tunes, or the set of inputs used to define the ID operating points for the ID. At this time, due to the success of the Likelihood used on offline data, it was also decided to be adapted for use in the trigger ID algorithms as well. I was involved in both the software changes and modifications to the method required by running in the online environment. In this chapter, I will briefly discuss electron reconstruction and the electron likelihood method.

4.1 Electron Reconstruction

Electron reconstruction is the process of creating electron candidates from the raw information collected by the triggering system. This is performed in several steps.

- **Seed-cluster reconstruction:** Energy deposits in the calorimeter are grouped into clusters. This is done with a sliding window algorithm summing the transverse energies in three layers of the EM calorimeter within an $\eta \times \phi$ region of $0.075 \times 0.125$.

- **Track reconstruction:** The path through the tracking volume is calculated creating a track.

- **Electron specific track refit:** Tracks are matched to a cluster. The $\eta$ and $\phi$ of the track are extrapolated into the second layer of the EM calorimeter. If this track is close enough to a cluster it is refit with a more advanced track fitter.

- **Electron candidate reconstruction:** The track matching of the previous step is redone with stricter criteria.

Because photons will look the same as electrons in the EM calorimeter, tracking criteria must be used to remove photons. Electron candidates that do not have a track associated to them are removed and considered photons. Electrons whose tracks are reconstructed to a
secondary vertex are also considered to be photons and are removed. Some electron candidates, based on their $E/p$, their $p_T$ and if there is an associated pixel hit can be classified as ambiguous, or unambiguous. This determines if it will be considered as only an electron, or if it will also be included in the photon container.

After all of these steps, some additional quantities are computed, such as the four-momentum, and the electron candidates have their energies calibrated.

4.2 Electron ID

4.2.1 Variables used for Electron ID

The various subsystems in the ATLAS detector can be used to make many measurements. The measurements used in the electron and photon ID are referred to as variables. These are described in table 4.1. A schematic drawing of the path of an electron through a section of the detector is depicted in figure 4.1.

The b-layer, silicon strips (SCT) and TRT all provide space point hits for tracking while the TRT also provides a number of high threshold hits that can be used to distinguish electrons from other particles that have similar detector signatures like pions. Tracking variables $d_0$ and $|d_0/\sigma_{d_0}|$ are measurements of the impact parameter, or the distance away from the beamline the particle originates from. These help to distinguish electrons from b-jets and c-jets that tend to have secondary vertices far from the beamline.

The TRT helps to discriminate electrons from heavier particles through the use of transition radiation. Lighter particles that have higher $\gamma$-factors will produce more photons when passing through detector material than heavier particles. This is measured with the variable $F_{HT}$ which is the ratio of high threshold hits to the total number of hits in the TRT. This tends toward higher values for electrons than for heavier particles.
4. ElectronID

Figure 4.1: A schematic drawing of a section of the ATLAS detector. The red line shows the path an electron takes through each subsystem [41].

The three main layers of the EM calorimeter provide shower shape information at different granularities. EM objects should not have showers longer than the EM calorimeter, so any energy deposited behind it into the hadronic calorimeter can be used to discriminate between electrons and light/heavy flavor objects. This is measured with the $R_{\text{had}}$ variable which is the ratio of transverse energy in the hardronic calorimeter to the EM calorimeter. The $f_1$ and $f_3$ variables measure the ratio of the energy in the first and third layers respectively to the total energy in the EM calorimeter. These allow transverse characteristics of the shower to be measured. The width the of the showers are measured with variables $w_{\text{stot}}$, $w_\eta$, $R_\eta$ and $R_\phi$. Electrons tend to have very narrow showers compared with heavier hadrons.

4.2.2 Electron Likelihood

The Electron Likelihood method developed to replace the cuts based ID is a naive Bayes classification method using probability density functions (PDFs) to determine the probability that a candidate is an electron. PDFs for both signal and background are created and are used
4. ElectronID

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>Name</th>
<th>Rejects</th>
<th>Use</th>
</tr>
</thead>
</table>
| Hadronic leakage         | Ratio of $E_T$ in the first layer of the hadronic calorimeter to $E_T$ of the EM cluster.  
(used over the range $|\eta| < 0.8$ or $|\eta| > 1.37$) | $R_{had1}$   | x       | x   | $\mathcal{L}$ |
|                          | Ratio of $E_T$ in the hadronic calorimeter to $E_T$ of the EM cluster.  
(used over the range $0.8 < |\eta| < 1.37$) | $R_{had}$    | x       | x   | $\mathcal{L}$ |
| Back layer of EM calorimeter | Ratio of the energy in the back layer to the total energy in the EM accordion calorimeter.  
This variable is only used for $p_T < 80$ GeV due to known inefficiencies at high $p_T$, and is also removed from the LH for $|\eta| > 2.37$, where it is poorly modeled by the MC. | $f_3$       | x       |      | $\mathcal{L}$ |
| Middle layer of EM calorimeter | Lateral shower width, $\sqrt{\sum (E_i \eta_i^2)/(\sum E_i) - (\sum E_i \eta_i)/(\sum E_i))^2}$, where $E_i$ is the energy and $\eta_i$ is the pseudorapidity of cell $i$ and the sum is calculated within a window of $3 \times 5$ cells | $w_\eta$    | x       | x   | $\mathcal{L}$ |
|                          | Ratio of the energy in $3 \times 3$ cells over the energy in $3 \times 7$ cells centered at the electron cluster position | $R_{\eta}$  | x       | x x | $\mathcal{L}$ |
|                          | Ratio of the energy in $3 \times 7$ cells over the energy in $7 \times 7$ cells centered at the electron cluster position | $R_{\eta}$  | x       | x x | $\mathcal{L}$ |
| Strip layer of EM calorimeter | Shower width, $\sqrt{\sum (E_i (i - i_{\text{max}})^2)/(\sum E_i)}$, where $i$ runs over all strips in a window of $\Delta \eta \times \Delta \phi \approx 0.0625 \times 0.2$, corresponding typically to 20 strips in $\eta$, and $i_{\text{max}}$ is the index of the highest-energy strip | $w_{\text{stot}}$ | x x x | G* |
|                          | Ratio of the energy difference between the maximum energy deposit and the energy deposit in a secondary maximum in the cluster to the sum of these energies | $\delta E_{\text{max}}$ | x       |      | $\mathcal{L}$ |
|                          | Ratio of the energy in the strip layer to the total energy in the EM accordion calorimeter | $f_1$       | x       |      | $\mathcal{L}$ |
| Track conditions         | Number of hits in the innermost pixel layer; discriminates against photon conversions | $n_{\text{Blayer}}$ | x       | C   |    |
|                          | Number of hits in the pixel detector | $n_{\text{Pixel}}$ | x       | C   |    |
|                          | Number of total hits in the pixel and SCT detectors | $n_{\text{Si}}$ | x       | C   |    |
|                          | Transverse impact parameter with respect to the beam-line | $d_{0}$     | x       | x   | $\mathcal{L}$ |
|                          | Significance of transverse impact parameter defined as the ratio of $d_{0}$ and its uncertainty | $|d_0/\sigma_{d_0}|$ | x       | x   | $\mathcal{L}$ |
| TRT                      | Likelihood probability based on transition radiation in the TRT | $e_{\text{ProbHT}}$ | x       |      | $\mathcal{L}$ |
| Track-cluster matching   | $\Delta \eta$ between the cluster position in the strip layer and the extrapolated track | $\delta_{\eta 1}$ | x x |      | $\mathcal{L}$ |
|                          | $\Delta \phi$ between the cluster position in the middle layer of the calorimeter and the momentum rescaled track extrapolated from the perigee | $\delta_{\phi_{e,x}}$ | x x |      | $\mathcal{L}$ |
|                          | Ratio of the cluster energy to the track momentum | $E/p$       | x       | x   | $\mathcal{C}^*$ |

Table 4.1: Definitions of electron discriminating variables, the types of backgrounds the variables help to discriminate against, and if a variable is used as a likelihood PDF ($\mathcal{L}$) or used as a rectangular cut (C). The * refers to the fact that the $E/p$ and $w_{\text{stot}}$ variables are only used for electrons with $p_T > 150$ GeV for the Tight identification operating point (in software release 20.7), and are not used for the looser operating points.
in combination to calculate a discriminant value. A cut is then placed on this discriminant to determine a desired signal efficiency or background rejection.

The likelihood is constructed by making a set of PDFs from a sample of electrons (signal) and electron fakes (background) for each of the \( n \) variables to be used. Each electron candidate to be classified has a set of the same \( n \) variables which we will call \( x \). The value from each signal (background) PDF associated with the variables \( x \) is calculated and their product is taken to get the signal (background) likelihood value. These are then combined to get the likelihood discriminant value according to equation (4.1).

\[
\begin{align*}
  d_L &= \frac{\mathcal{L}_S}{\mathcal{L}_S + \mathcal{L}_B}, \\
  \mathcal{L}_S(x) &= \prod_{i=1}^{n} P_{S,i}(x_i)
\end{align*}
\] (4.1)

This distribution for signal (background) tends to have a compressed peak at 1 (0) and so for finer control over the operating point a transformation is applied to the discriminant (4.2). This transformation broadens the peaks so the shape is more defined but still maintains signal and background separations.

\[
d_L' = -\tau^{-1} d_L \ln(d_L - 1), \quad \tau = 15
\] (4.2)

More advanced methods such as the electron likelihood have the advantage that they take into consideration multiple distributions at once, unlike the cuts method that looks at only one variable at a time. Figure 4.2 shows the signal and background distributions for the variables \( f_1 \) and \( R_\phi \). The large overlap of the signal and background causes any cut made on these variable to have poor background rejection. In some situations the variable might not be used at all as it adds no power to the method. With a likelihood, so long as there is a
difference in the shape, it can contribute to the power of the classification. Figure 4.3 shows the output discriminant of an example likelihood after transformation and the ROC curve corresponding to the continuum of operating points representing cuts on the discriminant. The good separation between signal and background can be seen.

Figure 4.2: Distribution of the variables $f_1$ and $R_\phi$ for signal (red) and background (blue). The large overlap of the signal and background of variables like this cause the cuts method to have poor background rejection.

Figure 4.3: An example likelihood discriminant output (a), after having been transformed by Equation 2, for data signal and background distributions. The corresponding ROC curve is shown in (b), illustrating the continuum of operating points. A cut-based menu is plotted for comparison.
A "tuned" likelihood consists of a set of PDFs, for each variable used in the likelihood and for each $\eta$ and $\phi$ bin used, and values of a cut on the discriminant for each desired operating point. The shape of the PDFs vary as a function of $p_T$ and $\eta$ so they are divided into $10 p_T$ and $9 \eta$ PDFs each with their own set of discriminant cuts. Tables 4.2 and 4.3 show the binning of the PDFs and discriminant cuts in $p_T$ and $\eta$. In addition to the discriminant cuts, one or more hits in the pixel tracker as well as seven or more SCT hits are required.

<table>
<thead>
<tr>
<th>Bin boundaries in $p_T$[GeV]</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDFs</td>
</tr>
<tr>
<td>4.5 7 10 15 20 30 40 $\infty$</td>
</tr>
<tr>
<td>Discriminant cut values</td>
</tr>
<tr>
<td>4.5 7 10 15 20 25 30 35 40 45 $\infty$</td>
</tr>
</tbody>
</table>

Table 4.2: Electron transverse energy binning used for the electron likelihood PDFs and discriminant cut values.

| Bin boundaries in $|\eta|$ |
|---------------------|
| 0.0 0.6 0.8 1.15 1.37 1.52 1.81 2.01 2.37 2.47 |

Table 4.3: Electron pseudorapidity binning used for the electron likelihood PDFs and discriminant cut values.

The discriminant values chosen are set to roughly match the efficiency of the 3 operating points (loose, medium, tight) used in the Run 1 offline likelihood. A fourth operating point was added (veryloose) with relaxed criteria compared to the loose operating point. This can be used to study backgrounds that are similar to signal electrons.

### 4.2.3 Probability Density Functions

PDFs can be obtained by making histograms of the variables used in the likelihood from signal and background samples. Finite statistics and the division of samples by $p_T$ and $\eta$ mean that the PDFs produced will not be smooth. This can lead to undesirable behavior in the likelihood as electrons in adjacent bins could have large differences in their respective
discriminants. In the tails of the PDF we also tend to have bins with 0 entries. In order to avoid undefined results, the PDFs should also be greater than zero everywhere. To correct this, we smooth the PDFs using an adaptive kernel density estimator (KDE). This takes each bin in the histogram and replaces it with a Gaussian distribution (the kernel) whose height and width are determined by the content of the bin. The difference between a non-adaptive and adaptive KDE is that Gaussian widths are variable in the adaptive case. This has the effect of narrowing the Gaussian kernel with increasing bin content as bins with high statistics have less uncertainty. The KDE technique is illustrated in figure 4.4 along with an example PDF before and after smoothing.

Figure 4.4: An example likelihood discriminant output (a), after having been transformed by Equation 2, for data signal and background distributions. The corresponding ROC curve is shown in (b), illustrating the continuum of operating points. A cut-based menu is plotted for comparison.

4.2.4 Correcting MC variable mis-modeling

One big advantage of the electron likelihood is that it can be tuned using data. A sample of electrons are gathered using the tag-and-probe method detailed in [42]. This sample will be a relatively unbiased, pure sample of data electrons with which to build PDFs. At the beginning
of Run 2, however, no data samples were available for the run conditions and center of mass energy the LHC was preparing for. This meant that we were forced to use MC to build the PDFs and tune the likelihood. These samples have some known mis-modeling effects in the form of shifts and in a few cases width scaling. This can be corrected however by comparing the MC variable distributions to $\sqrt{s} = 8$ TeV data and shifting or modifying the width of the distributions. Because of the difference in run conditions, the 8 TeV data will not exactly represent the variable shapes for 13 TeV data, but the shifts and widths will be close enough to use until data PDFs can be made with 13 TeV data.

The shifts are implemented as constant offsets of the mean position of the distributions. A comparison of data and MC showed that the variables which needed shifting corrections were $f_1$, $f_3$, $R_\eta$, $w_{\eta^2}$ and $R_\phi$. Width correction is performed by modifying the full width at half max (FWHM), which the variables $\Delta\eta_1$, $\Delta\phi_{\text{res}}$ and $R_{\text{had}}$ were found to need.

The operation to apply the shifts is performed on an electron-by-electron basis using $v_{\text{MC}}^* = v_{\text{MC}} - a$ for some offset $a$. To modify the widths the operation $v_{\text{MC}}^* = (v_{\text{MC}} - \bar{v}_{\text{MC}}) \cdot w + \bar{v}_{\text{MC}}$ for a width parameter $w$ and where $\bar{v}_{\text{MC}}$ is the mean of the distribution. The parameter $a$ is found by minimizing a $\chi^2$ test statistic,

$$\chi^2 = \sum_{\text{bins}} \frac{(n_{\text{data}} - n_{\text{MC}})^2}{\sigma_{\text{data}} + \sigma_{\text{MC}}}.$$  \hspace{1cm} (4.3)

The parameter $w$ is selected by taking the ratio of the FWHM of data to MC. Figure 4.5 shows the MC distributions before and after the corrections compared with the data distributions used.

While working in the Egamma group I was responsible for calculating all parameters of the shifts used to correct the MC. To make the corrections available to the entire group, I wrote a
tool that could be used to automatically correct all relevant distributions based on matrices of parameters that were kept updated in the tool. This was integrated into the official Egamma software package.

4.2.5 Signal and Background PDFs

The results of the KDE smoothing and MC correction procedure are shown in figures 4.6 and 4.7 for signal and background. The PDFs shown are taken from the $20 GeV < p_T < 30 GeV$, $0.6 < |\eta| < 0.8$ bin.

4.3 Pileup Correction

The LHC is continually trying to increase the luminosity of the collider. This allows more data to be collected in a shorter amount of time. The increase in luminosity means that the average number of interactions per bunch crossing also increases. In 2016 the average was $\sim 25$ interaction, up from $\sim 12.5$ in 2015. The additional vertices in each event contribute to what is known as pileup. Particles from completely different interaction vertices can overlap, depositing energy in the same regions of the detector. Some ID variables are particularly effected by this and their shapes, in general, become broader. $R_\eta$ and $R_{had}$ have the strongest dependence on pileup as seen in figure 4.8. When these distributions become wider, their PDFs become more background like. For events with a lot of pileup, this causes a drop in the efficiency to identify electrons. $R_\eta$ and $R_{had}$ are among the variables with the highest discriminating power as can be seen in figure 4.9 and so they cannot just be removed from the likelihood.

We would like to correct for this drop in efficiency. We can do this by making the discriminant cut linearly dependent on the number of primary vertices ($n_{vtx}$), $d(n_{vtx}) = d_L + a n_{vtx}$. 
Figure 4.5: Data and MC electron variable distributions obtained using the $Z \to ee$ tag-and-probe method, and in the $30 \leq E_T \leq 40$, $0.80 \leq \eta \leq 1.15$ bin. MC is shown before and after applying the constant shift and width corrections described in the text. KDE smoothing has been applied.
Figure 4.6: MC-based signal and background PDFs used in the electron LH for software release 20.7 (with MC shower shape shifts and widths applied), shown for the variables $f_1$, $f_3$, $R_\eta$, $R_\phi$, $R_{had}$, $E_{ratio}$, and $w_\eta$, after KDE smoothing has been applied. Only electron candidates with $20 \text{ GeV} < E_T < 30 \text{ GeV}$ and $0.60 < |\eta| < 0.80$ are shown. As the MC was used to construct these PDFs, the probes from $Z \rightarrow ee$ were additionally required to pass a truth matching requirement, while the background electron candidates from a dijet sample were required to fail the electron truth matching.
Figure 4.7: MC-based signal and background PDFs used in the electron LH for software release 20.7 (with MC shower shape shifts and widths applied), shown for the variables eProbabilityHT, $d_0$, $|d_0/|d_0||$, $\Delta \eta_1$, $\Delta \phi_{res}$, and $\Delta p/p$, after KDE smoothing has been applied. Only electron candidates with $20 \text{ GeV} < E_T < 30 \text{ GeV}$ and $0.60 < |\eta| < 0.80$ are shown. As the MC was used to construct these PDFs, the probes from $Z \rightarrow ee$ were additionally required to pass a truth matching requirement, while the background electron candidates from a dijet sample were required to fail the electron truth matching.
Figure 4.8: \( R_\eta \) is shown in (a) integrated for Nvtx < 11 and Nvtx > 11. The same is shown for \( R_{had} \) in (b).

Figure 4.9: The \( n - 1 \) method used to optimize the choice of variables to use in the electron likelihood. Individual variables are removed from the nominal list of likelihood variables, and the likelihood recalculated to assess the relative power of each variable. The example shows the importance of \( F_{HT} \), \( E_{ratio} \), \( R_{had} \) and \( R_\eta \); the performance of the likelihood decreases when each is removed. The Tight cut-based operating point is shown for comparison.
This can correct for the negative slope in the signal efficiency, but because we are relaxing the cut at high \( n_{vtx} \) we are also letting more background pass the cut. This situation is illustrated in figure 4.10.

![Figure 4.10: In black are the efficiencies corresponding to the signal efficiency tuning being linearly corrected as a function of \( n_{vtx} \). We observe how the background blows up for this first order correction. In red we see the second correction in where the signal efficiency is re-injected with some slope in order to stabilize the background.](image)

This correction as a function of \( n_{vtx} \) is applied to only the Medium and Tight operating points as the pileup dependence for Loose and VeryLoose are too small to have a noticeable effect. In the plane defined by the likelihood discriminant as a function of \( n_{vtx} \) this correction would correspond to tilting the line defining the cut giving it non-zero slope. However, since the Loose and VeryLoose operating points are not adjusted, there are situations where at high pileup the line defining the Medium operating point will cross the looser ones. The electrons passing the Medium likelihood is then no longer strictly a subset of the Loose likelihood. Figure 4.11 shows how this situation can arise.

The remedy for this situation we decided to make was to keep the discriminant cuts flat as a function of \( n_{vtx} \) and instead correct the discriminant output as a function of the pileup. This will ensure that all tighter operating points are subsets of looser ones by construction.
Figure 4.11: Example of the pileup correction used in Run 1 where the z-axis shows the number of entries in each bin. The blue dashed line shows that with large pileup, the Medium operating point could be looser than the Loose operating point, so they are not perfect subsets. The solid blue line shows what is desired, namely, for Medium to always be a subset of Loose.

The correction to the discriminant is done in the following way:

\[
d_{\text{new}}(n_{\text{vtx}}) = \begin{cases} 
  d & \text{if } d < d_{\text{ref}} \\
  d_{\text{ref}} + (d - d_{\text{ref}})\frac{d_{\text{tight,old}} - d_{\text{ref}}}{d_{\text{tight,new}} - d_{\text{ref}}} & \text{if } d_{\text{ref}} \leq d < d_{\text{tight,new}} \\
  d_{\text{tight,old}} + (d - d_{\text{tight,new}})\frac{d_{\text{max}} - d_{\text{tight,new}}}{d_{\text{max}} - d_{\text{tight,new}}} & \text{if } d_{\text{tight,new}} \leq d < d_{\text{max}} \\
  d & \text{if } d_{\text{max}} \leq d 
\end{cases} 
\]  

(4.4)

Here \(d_{\text{ref}}\) is a reference discriminant which defines the lowest point where pileup corrections are to be done. This is set to the VeryLoose operating point as we do not require it to be pileup corrected. Similarly \(d_{\text{max}}\) is the largest value of the discriminant that should be corrected.

This method was found to have performance similar to the Run 1 scheme of correcting the cut values themselves, but ensures that the tighter points remain subset of the looser.
4.3.1 Conclusion

The electron likelihood used offline in Run 1 has been extremely successful. Its improvements over the cuts based ID made it a good candidate for operation in the trigger in Run 2 where the demands on the ID would be higher. The operating points implemented for online and offline running closely match the signal efficiencies of the cuts based ID, but with better background rejection. Although the luminosity of the LHC is steadily increasing, the likelihood is able to adjust around this and will continue to deliver steady performance through the end of Run 2.
CHAPTER 5

The Search for Low Mass Diphoton Resonances

Most of the activities on Atlas can be divided into measurements and searches. Measurement groups find a numerical value for some parameter of a previously discovered particle or process. An example of this would be the Higgs cross section group which makes ever more accurate measurements of the cross section and branching ratios of the Higgs Boson. In order to make these measurements, the particle whose parameters you will be assessing need to be discovered. This is where the search groups come in. Search groups have the goal of finding new particles and processes. These measurements are simply to determine if there is a statistically significant deviation from currently known physics. Where I spent a majority of my time on Atlas was in a group interested in searching for new particles that decay to two photons and appear in the data as resonances, or narrow bumps.

In the following I will present a search for low mass diphoton resonances during 2017 and 2018 [43]. This analysis used $pp$ collision data with a total integrated luminosity of $80.4 \ fb^{-1}$ and a center of mass energy of $\sqrt{s} = 13\text{TeV}$ collected in 2015, 2016 and 2017. This search is an update of the Run 1 results with $20.3\ fb^{-1}$ of data at $\sqrt{s} = 8\text{TeV}$ [44] The search is performed for narrow resonances with spin 0. Experimental signatures such as these are predicted by models involving an extended Higgs sector and some Supersymmetric/Axion models.
This analysis represents the work of many people over many months. This chapter closely follows the low mass diphoton support note [43] meant to supplement the conference note [45] both of which I spent the last year or so working on. Although I was responsible for many support and production tasks such as data validation, and signal Monte Carlo production, my main focus was on the non-resonant background modeling and the spurious signal test described in section 5.6.3.

5.1 Introduction

Several theoretical models predict new resonances in the diphoton invariant mass spectrum. One of the more prominent are extensions to the Higgs sector of the standard model. The Standard Model Higgs sector contains one complex scalar doublet field that gives rise to the standard model Higgs boson. This is the minimal scalar sector that can be added to the Standard Model to give mass to particles through the Higgs mechanism. The scalar sector does not necessarily need to be minimal however.

The simplest extension is to add one additional complex scalar doublet field. This is known as the two Higgs doublet model (2HDM) [46] [47] [48]. The scalar potential for two doublets can be written,

\[
V = m_{11}^2 \Phi_1^\dagger \Phi_1 + m_{22}^2 \Phi_2^\dagger \Phi_2 - m_{12}^2(\Phi_1^\dagger \Phi_2 + \Phi_2^\dagger \Phi_1) + \frac{\lambda_1}{2} (\Phi_1^\dagger \Phi_1)^2 + \frac{\lambda_2}{2} (\Phi_2^\dagger \Phi_2)^2 \\
+ \lambda_3 \Phi_1^\dagger \Phi_1 \Phi_2^\dagger \Phi_2 + \lambda_4 \Phi_1^\dagger \Phi_2^\dagger \Phi_1 + \lambda_5 \frac{1}{2} \left[ (\Phi_1^\dagger \Phi_2)^2 + (\Phi_2^\dagger \Phi_1)^2 \right].
\]

The minimization of this potential is,

\[
\langle \Phi_1 \rangle_0 = \begin{pmatrix} 0 \\ \frac{v_1}{\sqrt{2}} \end{pmatrix}, \quad \langle \Phi_2 \rangle_0 = \begin{pmatrix} 0 \\ \frac{v_2}{\sqrt{2}} \end{pmatrix}.
\]

With two complex scalar doublets, there will be 8 fields in total. As in the SM Higgs, 3 of these fields get "eaten" to give mass to the $W^\pm$ and $Z$ bosons. This leaves 5 physical scalar
fields; two charged scalars, two neutral scalars, and one pseudoscalar.

The Yukawa couplings in the 2HDM [47] in general, will be,

\[ \mathcal{L}_y = y_{ij}^1 \bar{\psi}_i \psi_j \Phi_1 + y_{ij}^2 \bar{\psi}_i \psi_j \Phi_2, \]  

(5.2)

with mass matrix,

\[ M_{ij} = y_{ij}^1 \frac{v_1}{\sqrt{2}} + y_{ij}^2 \frac{v_2}{\sqrt{2}} \]  

(5.3)

In the Standard Model, the mass matrix and Yukawa matrices diagonalize meaning that there are no flavor changing neutral currents (FCNC). With the addition of an additional scalar doublet, the Yukawa matrix is not necessarily guaranteed to be diagonalized if the mass matrix is. This means that the 2HDM can have FCNC at tree level which have very strong constraints from experiment [49]. To avoid this problem an additional \( Z_2 \) symmetry can be enforced. This can be applied in a few different ways; the type I where all fermions only couple to one of the doublets, and the type II where the up-type quarks couple to one doublet, with the down-type quarks and leptons coupling to the other. Two other models are possible (type III and type IV) which do allow tree level FCNC but constrain them in other ways.

The two neutral scalars are \( h \) and \( H \) with \( h \) the lighter of the two. Either one of these two could be the SM Higgs and so the second neutral scalar could be found above or below 125 GeV. Just like in the SM the branching ratio of bosons to photons in the 2HDM are not predicted to be large, but the clean experimental signature and high resolution of the ATLAS detector make the diphoton channel very good at searching for an extended scalar sector.
5. Low Mass Diphoton

5.2 Data and Simulation

5.2.1 Data

In 2015 the ATLAS experiment began the Run 2 data collection period where the center of mass energy was increased from \( \sqrt{s} = 8\) TeV to near its design energy of \( \sqrt{s} = 13\) TeV. For this analysis, data from 2015, 2016 and 2017 were used. Events were recorded by requiring a diphoton trigger with a transverse energy \( (E_T) \) of 20 GeV on both the leading and subleading photon candidates. The high level trigger (HLT) then requires that both photons pass tight reconstruction criteria in addition to being isolated for the 2017 data only. The data used after these requirements corresponds to \( 80fb^{-1} \) of total integrated luminosity [43].

5.2.2 Signal Simulation

Monte Carlo simulation (MC) is used to produce both signal and background samples that are used for various modeling tasks as well as to optimize the search strategy. Due to the difference in run conditions between the 2015/2016 and the 2017 data collection periods, simulation must be generated separately for each. The mc16a samples correspond to the 2015/2016 and the mc16d samples to the 2017 data.

The different MC signal samples are detailed in Table 5.1. The default samples are the ggF ones, generated using the effective-field-theory approach implemented in MadGraph5_aMC@NLO version 2.3.3 at next-to-leading order (NLO) in quantum chromodynamics (QCD). From the Higgs characterization framework [51], CP-even dimension-five operators coupling the new resonance to gluons and photons were included. Samples were generated with the NNPDF3.0 NLO parton distribution functions (PDFs) [52], using the A14 set of tuned parameters (tune) of Pythia 8.186 [53, 54] for the parton-shower and hadronization simulation. Narrow Width Approximation is assumed for all samples (a 4 MeV width is used in practice). Interference
effects between the resonant signal and the background processes are neglected. Mass points outside of the search region (65-110 GeV) were generated in order to have a better interpolation of the signal modeling (cf Section 5.4). But in the end the 40 and 50 GeV mass points are not used because the shape is biased by the trigger cut.

Since the goal of the analysis is to perform a model-independent search, the properties of the fiducial volume and correction factors (detailed in Section 5.8) are also studied for other Higgs-like production modes: vector boson fusion (VBF), associated production with a vector boson (WH, ZH) and associated production with a top quark pair ($t\bar{t}H$). All those samples are simulated with the PowHeg\textsuperscript{55, 56} generator both interfaced to Pythia8\textsuperscript{53} for parton showering and hadronization.

<table>
<thead>
<tr>
<th>Process</th>
<th>Generator</th>
<th>Mass [GeV]</th>
<th>Nevents ($\times 10^3$)</th>
<th>width</th>
</tr>
</thead>
<tbody>
<tr>
<td>ggF</td>
<td>MadGraph</td>
<td>40-50-60-70-80-90-100-110-120-140-160-180</td>
<td>mc16a 30</td>
<td>5%</td>
</tr>
<tr>
<td>ggF</td>
<td>PowHeg+Pythia8</td>
<td>40-60-80-100-120</td>
<td>mc16d 30</td>
<td>NWA</td>
</tr>
<tr>
<td>VBF</td>
<td>PowHeg+Pythia8</td>
<td>40-60-80-100-120</td>
<td>30</td>
<td>NWA</td>
</tr>
<tr>
<td>WH</td>
<td>Pythia8</td>
<td>40-60-80-100-120</td>
<td>30</td>
<td>NWA</td>
</tr>
<tr>
<td>ZH</td>
<td>Pythia8</td>
<td>40-60-80-100-120</td>
<td>30</td>
<td>NWA</td>
</tr>
<tr>
<td>$t\bar{t}H$</td>
<td>PowHeg+Pythia8</td>
<td>40-60-80-100-120</td>
<td>30</td>
<td>NWA</td>
</tr>
</tbody>
</table>

Table 5.1: Signal samples of the five Higgs production modes, available mass points and whether the NWA width is used or not 43.

5.2.3 Background Simulation

MC samples of prompt diphoton events are used to validate the functional form used to model the background diphoton invariant mass spectrum.

Events containing two photons are also generated using the Sherpa 2.2.2 generator, with a 20 GeV cut on the photon transverse energy. Matrix elements are calculated with up to 2 partons at LO and merged with the Sherpa parton shower 57 using the ME+PSatLO
5. Low Mass Diphoton

<table>
<thead>
<tr>
<th>Generator</th>
<th>$m_{\gamma\gamma}$ range [GeV]</th>
<th>Cross section [pb]</th>
<th>Filter efficiency</th>
<th>Nevents ($\times 10^6$)</th>
<th>Reconstruction brawl 16a brawl 16d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sherpa LO</td>
<td>0 – 55</td>
<td>$1.4088 \times 10^{+2}$</td>
<td>$2.4335 \times 10^{-1}$</td>
<td>1.0</td>
<td>1.0 FS</td>
</tr>
<tr>
<td></td>
<td>55 – 100</td>
<td>$1.4778 \times 10^{+2}$</td>
<td>$4.5670 \times 10^{-1}$</td>
<td>1.0</td>
<td>1.0 FS</td>
</tr>
<tr>
<td></td>
<td>100 – 160</td>
<td>$3.9728 \times 10^{+1}$</td>
<td>$4.9730 \times 10^{-1}$</td>
<td>1.0</td>
<td>1.0 FS</td>
</tr>
<tr>
<td>Sherpa NLO</td>
<td>50 – 90</td>
<td>$1.3904 \times 10^{+2}$</td>
<td>1</td>
<td>91.7</td>
<td>130.0 AFII</td>
</tr>
<tr>
<td></td>
<td>90 – 175</td>
<td>$5.1823 \times 10^{+1}$</td>
<td>1</td>
<td>114.7</td>
<td>162.7 AFII</td>
</tr>
</tbody>
</table>

Table 5.2: Prompt diphoton background samples.

prescription [58]. The CT10 PDF set is used in conjunction with dedicated parton shower tuning developed by the Sherpa authors. Those samples are used for optimisation of the analysis and for cross-checks.

A recent implementation of Sherpa, version 2.2.4 [59], is also used. It consistently combines parton-level calculations of varying jet multiplicity up to NLO (The $\gamma\gamma$ and $\gamma\gamma+1$ parton processes are generated at NLO accuracy, while the $\gamma\gamma+2$ partons and $\gamma\gamma+3$ partons processes are generated at LO. Charm and bottom quarks are included in these matrix elements in the massless approximation.) with parton showering while avoiding double-counting effects. The NNPDF3.0 NNLO PDFs [60] are used in conjunction with the corresponding Sherpa default tuning. Those samples with high-statistics are used for the background modelling studies described in Section 5.5.

Details on those samples are given in Table 5.2. In order to maximize the available statistics over the mass range of interest, the simulation is separately performed in exclusive slices of diphoton invariant mass. Those samples are fully reconstructed and are used for optimization of the analysis.

Samples with one photon and up to 2 jets are simulated with Sherpa and are generated in slices of the photon transverse energy. They are described in the general HGam supporting note [61] and are used for the study of the isolation cuts.

Monte-Carlo samples of $Z \rightarrow ee$ decays are simulated using POWHEG-BOX V2 interfaced
5. Low Mass Diphoton

<table>
<thead>
<tr>
<th>Generator</th>
<th>( m_{ee} ) range [GeV]</th>
<th>Filter</th>
<th>Cross section [pb]</th>
<th>Filter efficiency</th>
<th>Nevents ((\times 10^6))</th>
</tr>
</thead>
<tbody>
<tr>
<td>POWHEG</td>
<td>–</td>
<td>c-veto, b-veto</td>
<td>1.9810 \times 10^{-3}</td>
<td>0.82143</td>
<td>8.0</td>
</tr>
<tr>
<td></td>
<td>0 – 70</td>
<td>c-filter, b-veto</td>
<td>1.9816 \times 10^{-3}</td>
<td>0.11407</td>
<td>5.0</td>
</tr>
<tr>
<td></td>
<td>70 – 140</td>
<td>c-veto, b-veto</td>
<td>1.1063 \times 10^{-3}</td>
<td>0.69432</td>
<td>6.0</td>
</tr>
<tr>
<td></td>
<td>70 – 140</td>
<td>c-filter, b-veto</td>
<td>1.1045 \times 10^{-2}</td>
<td>0.18697</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td>140 – 280</td>
<td>c-veto, b-veto</td>
<td>4.0711 \times 10^{-1}</td>
<td>0.61632</td>
<td>5.0</td>
</tr>
<tr>
<td></td>
<td>140 – 280</td>
<td>c-filter, b-veto</td>
<td>4.0683 \times 10^{-1}</td>
<td>0.23302</td>
<td>3.0</td>
</tr>
<tr>
<td></td>
<td>280 – 500</td>
<td>c-veto, b-veto</td>
<td>8.6711 \times 10^{+0}</td>
<td>0.56328</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td>280 – 500</td>
<td>c-filter, b-veto</td>
<td>8.6597 \times 10^{+0}</td>
<td>0.26640</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td>280 – 500</td>
<td>b-filter</td>
<td>8.6793 \times 10^{+0}</td>
<td>0.17638</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td>500 – 1000</td>
<td>c-veto, b-veto</td>
<td>1.8096 \times 10^{+0}</td>
<td>1</td>
<td>3.0</td>
</tr>
<tr>
<td></td>
<td>1000 – \infty</td>
<td>–</td>
<td>1.4875 \times 10^{-1}</td>
<td>1</td>
<td>1.0</td>
</tr>
</tbody>
</table>

Table 5.3: \( Z \to ee \) samples used to study the Drell-Yan background.

to the PYTHIA8 version 8.186 parton shower model. The CT10 parton density function set is used in the matrix element. The AZNLO set of tuned parameters is used, with PDF set CTEQ6L1, for the modeling of non-perturbative effects. Sliced samples generated with Sherpa 2.2.1 are also used for computing systematic uncertainties. Details on those samples are given in Table 5.3.

The data and MC samples are processed through the HGam framework, and the mini-xAODs version h021 are used unless stated otherwise. Weights are applied to correct the simulation: Scale Factor for the photon identification, pile-up, choice of the diphoton vertex, and MC weight when applicable (eg Sherpa samples).
5. Low Mass Diphoton

5.3 Event Selection

5.3.1 Trigger Selection

In general we want to be able to perform this search as low as possible in diphoton mass. One of the limits we run into in this regard is the threshold at which triggers become prescaled. At very low transverse energies \( E_T \), the rate that events occur in the detector is much higher than the detector is able to record at. This means that low \( E_T \), the triggers must be prescaled, or have their rate limited. We would like to use triggers that are unprescaled as they will collect a higher fraction of the events produced above their threshold. For 2015 and part of 2016 running the trigger 2g20_tight was used, which corresponds to a diphoton trigger where both the leading and subleading photons are required to have \( E_T > 20 \) GeV and both pass tight identification. For the later part of 2016 running, the threshold was raised to 22 GeV for the lowest unprescaled trigger (2g22_tight) due to the increase in luminosity of the LHC. In 2017, the threshold was lowered back to 20 GeV by adding the additional requirement that the photons pass the very loose calorimeter isolation (2g20_tight_icalovloose). The triggers used are summarized in table 5.4.

<table>
<thead>
<tr>
<th>Year</th>
<th>2015</th>
<th>2016 up to D3</th>
<th>2016 from D3</th>
<th>2017</th>
</tr>
</thead>
<tbody>
<tr>
<td>HLT item</td>
<td>2g20_tight</td>
<td>2g20_tight</td>
<td>2g22_tight</td>
<td>2g20_tight_icalovloose</td>
</tr>
<tr>
<td>luminosity [fb⁻¹]</td>
<td>3.2 fb⁻¹</td>
<td>11.5</td>
<td>21.5</td>
<td>43.6 fb⁻¹</td>
</tr>
</tbody>
</table>

Table 5.4: Lowest unprescaled HTL items, depending on the data-taking period, and associated integrated luminosity [43].

The use of 20 GeV triggers means we can use an offline cut of 22 GeV on the photon \( E_T \). This causes a kinematic turn on in the invariant mass distribution which peaks at around 60 GeV. Although this turn on makes background modeling more difficult (see section 5.6.3), it allows us to fit the distribution down to 60 GeV. The efficiency of the triggers are studied by
the Egamma group’s photon trigger experts and are shown in figure 5.1a. These efficiencies have good agreement with simulation and are shown in figure 5.1b as the ratio of data to MC.

Figure 5.1: \texttt{g20\_tight\_icalovloose} trigger efficiencies (a) and data/MC ratios (b) measured on 2017 data.
5. Low Mass Diphoton

5.3.2 Kinematic Selection

As mentioned before we use an offline cut of \( E_T > 22 \text{ GeV} \). We want to keep this as low as possible for two reasons: to be able to start the search as low as possible and to keep the kinematic turn on caused by the cut far away from the Drell-Yan peak. Changes to this cut were not considered.

Also considered were relative cuts where the absolute \( E_T \) cut is also divided by the invariant mass of the pair of photon candidates. These relative cuts are used in the SM \( H \rightarrow \gamma\gamma \) analysis [62] as well as the high-mass diphoton analysis [8]. Figure 5.2a shows a comparison of the absolute and relative cuts defined as \( E_T > 22\text{GeV/}X\text{GeV} \). You can see that for the relative cuts the kinematic turn on follows the absolute cut until the value of \( X\text{GeV} \) where it falls off. To measure the effect on the background we use background only SHERPA MC samples described in section 5.2. The number of background events are measured in a window centered at \( m_X \), the mass of the signal point, with a width of \( 2 \times \sigma_{CB} \) where \( \sigma_{CB} \) is the width of the double sided crystal ball function used in the signal parameterization described in section 5.4. Figure 5.2b shows the comparison of the tested cuts on the number of background events normalized to the \( E_T > 22\text{GeV} \) cut. The same is done in figure 5.2c for the number of signal events measured on the ggF signal MC samples, again normalized to the absolute cut.

In order to test the cuts, we use the significance \( Z = S/\sqrt{B} \). This analysis is process independent and so we have no expected number of events to compare. The ratio of \( S/\sqrt{B} \) for the relative cut divided by \( S/\sqrt{B}_{E_T > 22\text{GeV}} \) for the default 22GeV cut is shown in figure 5.2d. The significance is much lower at high masses for the relative cuts so it was concluded that the default absolute cut \( E_T > 22\text{GeV} \) will be used for this analysis.
Figure 5.2: (a) $M_{\gamma\gamma}$ distribution of the diphoton background for various sets of kinematic cuts applied on the photon transverse energies. (b) Ratio between the number of background events in $m_X \pm 2 \times \sigma_{CB}$ with the tested cuts and with the 22 GeV cuts. (c) Ratio between the number of signal events in $m_X \pm 2 \times \sigma_{CB}$ with the tested cuts and with the 22 GeV cuts. (d) Ratio between $S/\sqrt{B}$ with the tested cuts and $S/\sqrt{B}$ with the 22 GeV cuts.
5.3.3 Isolation Selection

In addition to using an isolation requirement on the 2017 trigger, we also use isolation in the offline selection of events. Isolation ensures that there is not too much activity in the detector around the photon from both other interaction vertices but also from other decay product originating from the primary vertex. Non-isolated candidates have a higher probability of being misconstructed as photons. Two types of isolation variables are used; track-based and calorimeter-based. Track-based isolation, $ptcone_{20}$, is defined as the scalar sum of the transverse momenta of all tracks with $p_T > 1\text{GeV}$ in the cone of size $\Delta R = \sqrt{(\Delta \eta)^2 + (\Delta \phi)^2} = 0.2$ around each photon candidate. Only tracks that are reconstructed as originating from the primary vertex are used, and tracks associated with conversions are also removed. The calorimeter-based isolation uses only the topological clusters in the calorimeter. $topoetcone_{40}$ is the sum of all positive energy in clusters within a cone of $\Delta R = 0.4$ after having subtracted the contribution from the photon candidate itself.

Similar to how the photon ID is done, there are three different isolation cuts that are available as working points:

- **FixedCutTightCaloOnly**: $topoetcone_{40} < 0.022p_T + 2.45\ [\text{GeV}]$, aiming for SM measurements;

- **FixedCutTight**: $topoetcone_{40} < 0.022p_T + 2.45\ [\text{GeV}]$ and $ptcone_{20}/p_T < 0.05$, aiming for high-$E_T$ photons;

- **FixedCutLoose**: $topoetcone_{20} < 0.065p_T$ and $ptcone_{20}/p_T < 0.05$, designed for the SM $H \rightarrow \gamma\gamma$ analysis.

In order to choose which working point to use, we look at the significance for each cut. We will measure the significance relative to the FixedCutLoose working point by taking the ratio
### Table 5.5: $m_{\gamma\gamma}$ cuts applied on signal and background samples with respect to the different mass of resonance points.

<table>
<thead>
<tr>
<th>Generator of signal samples</th>
<th>Mass point [GeV]</th>
</tr>
</thead>
<tbody>
<tr>
<td>MadGraph</td>
<td>$M_X = 40, 50, 60, 70, 80, 90, 100, 110$</td>
</tr>
<tr>
<td></td>
<td>$M_X - 5 &lt; m_{\gamma\gamma} &lt; M_X + 5 \text{ GeV}$</td>
</tr>
<tr>
<td>PowHeg+Pythia8</td>
<td>$M_X = 40, 60, 80, 100, 120$</td>
</tr>
<tr>
<td></td>
<td>$M_X - 10 &lt; m_{\gamma\gamma} &lt; M_X + 10 \text{ GeV}$</td>
</tr>
</tbody>
</table>

of the two. This gives us the relative efficiency $\epsilon_{\text{relative}} = \frac{N_{\text{testcut}}}{N_{\text{FixedCutLoose}}}$ where $N_{\text{testcut}}$ ($N_{\text{FixedCutLoose}}$) is the number of events passing the cut we are testing (number of events passing FixedCutLoose). The gain or loss in significance as measured relative to FixedCutLoose is written as $Z_{\text{testCut}}/Z_{\text{FixedCutLoose}} = \epsilon_S/\sqrt{\epsilon_B}$ where $\epsilon_S$ and $\epsilon_B$ are the relative efficiencies for the signal and background respectively.

The samples primarily used for the isolation study are the MadGraph samples, however, samples produced with PowHeg+Pythia8 are used as well to check the effect of the working points on different production modes. These samples are detailed in section 5.2.2. For the background samples we use the Sherpa leading order samples in addition to the Sherpa $\gamma$ jet samples. The $\gamma$ jet component ranges from about 50% to 30% of the total background events according to the study in section 5.6.1. This means we need to account for this contribution as jets have a different isolation distribution from photons. Isolation working points are tested for two fraction of $\gamma$ jet contribution, 50% $\gamma\gamma$: 50% $\gamma$ jet and 70% $\gamma\gamma$: 30% $\gamma$ jet. Jet-jet components are neglected as they are small ($\sim 6\%$ section 5.6.1).

The events that are used for testing are required to pass the basic kinematic selection of $2g20_{\text{tight}}$ trigger, tight identification and $E_T > 22 \text{ GeV}$. There is then an addition cut placed on the events invariant mass in a window around the mass of the signal being tested. These cuts range from ±5GeV for lower masses to ±20 GeV for higher masses in the PowHeg samples. These $M_{\gamma\gamma}$ cuts are shown in table 5.5.
Figure 5.3 shows the relative isolation efficiencies for the background samples. The binning in this plot corresponds to the mass points in the MadGraph signal sample so that they may be easily compared and combined. Likewise, figure 5.4 shows the same background samples, but with the binning corresponding to the mass points in the PowHeg signal samples. As can be seen, the ratios for the FixedCutTight isolation are below 1, showing that this working point has a higher rejection than the FixedCutLoose. This is expected as the tight cuts are designed to have a greater rejection (or lower efficiency). The relative efficiency for the FixedCutTightCaloOnly is greater than or close to 1, showing that its rejection power is lower than that of the track-based FixedCutLoose working point.

The relative efficiency for the MadGraph signal samples are shown in figure 5.5. We find a similar situation to the background efficiency with the FixedCutTightCaloOnly being more efficient than FixedCutLoose, and FixedCutTight being slightly less. Figure 5.6 shows the significance relative to the FixedCutLoose working point for the two different fractions of $\gamma$-jet. In both cases, the FixedCutTight working point has a higher significance than the calorimeter only version, but both are lower and 1 in general. Although the FixedCutTight isolation is very close to one for a few mass points, it is on average lower.

Figure 5.7 shows the relative efficiencies for the PowHeq + Pythia8 signal samples with the ggH, ttH, VBF, WH, and ZH production modes. The significance for each of these production modes is shown in figure 5.8 for the 70% : 30% $\gamma$-jet fraction, and the same in figure 5.9 for the 50% : 50% $\gamma$-jet fraction. We see similar results for the PowHeg + Pythia8 samples as for the MadGraph samples. The FixedCutLoose working point tends to have the highest significance across the invariant mass range and so it is used for the analysis.
Figure 5.3: Isolation efficiency relative to the FixedCutLoose working point for background samples: (a) $\gamma\gamma$. (b) $\gamma\text{jet}$. (c) combined background sample with $\gamma\gamma : \gamma\text{jet} = 70\% : 30\%$. (d) $\gamma\gamma : \gamma\text{jet} = 50\% : 50\%$. Efficiencies for the FixedCutTight (respectively FixedCutTightCaloOnly) working point are represented with full (respectively open) circles.
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Figure 5.4: Isolation efficiency relative to the FixedCutLoose working point for background samples: (a) $\gamma\gamma$. (b) $\gamma\text{jet}$. (c) combined background sample with $\gamma\gamma : \gamma\text{jet} = 70% : 30%$. (d) $\gamma\gamma : \gamma\text{jet} = 50% : 50%$. Efficiencies for the FixedCutTight (respectively FixedCutTightCaloOnly) working point are represented with full (respectively open) circles.
Figure 5.5: Isolation efficiency relative to the FixedCutLoose working point for MadGraph signal samples. Black open circles represents that FixedCutTight isolation requirement is applied, and blue dots represents FixedCutTightCaloOnly.

Figure 5.6: Significance relative to the FixedCutLoose working point with MadGraph signal sample and background contains: (a) $\gamma\gamma : \gamma jet = 70\% : 30\%$. (b) $\gamma\gamma : \gamma jet = 50\% : 50\%$. Significances for the FixedCutTight (respectively FixedCutTightCaloOnly) working point are represented with full (respectively open) circles.
Figure 5.7: Isolation efficiency relative to the FixedCutLoose working point for signal samples:
(a) ggH. (b) ttH. (c) VBFH. (d) WH. (e) ZH. Efficiencies for the FixedCutTight (respectively
FixedCutTightCaloOnly) working point are represented with full (respectively open) circles.
Figure 5.8: Significance relative to the FixedCutLoose working point with signal sample as: (a) ggH. (b) ttH. (c) VBFH. (d) WH. (e) ZH. The background contains $\gamma\gamma : \gamma jet = 70\% : 30\%$. Significances for the FixedCutTight (respectively FixedCutTightCaloOnly) working point are represented with full (respectively open) circles.
Figure 5.9: Significance relative to the FixedCutLoose working point with signal sample as: (a) ggH. (b) ttH. (c) VBFH. (d) WH. (e) ZH. The background contains $\gamma\gamma : \gamma{jet} = 50\% : 50\%$. Significances for the FixedCutTight (respectively FixedCutTightCaloOnly) working point are represented with full (respectively open) circles.
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5.3.4 Categorization

The determination of which and how many categories to use is important for this analysis as we expect to have different numbers of Drell-Yan events for different conversion status of the photon candidates. Conversions happen when photons in material convert into $e^+e^-$ pairs. Conversion status is split into 3 categories. Unconverted, converted with 1 track, and converted with 2 tracks. The two conversion categories are then further subdivided by which detector sub systems (silicon tracker or TRT) the tracks appear in.

- **unconverted**: unconverted photon

- **conv 1-track**: single track converted photon, divided in:
  - singleSi: one track only, with Si hits
  - singleTRT: one track only, no Si hits (TRT only)

- **conv 2-track**: double track converted photon, divided in:
  - doubleSi: two tracks, both with Si hits
  - doubleTRT: two tracks, none with Si hits (TRT only)
  - doubleSiTRT = two tracks, only one with Si hits

Three different categorization schemes are considered where the leading and subleading photons conversion status are treated separately.

- **4 Categories** - Checking only whether the photon is converted or not for leading and subleading.

- **9 Categories** - As above but splitting the conversions into 1-track and 2-track varieties.
5. Low Mass Diphoton

- 36 Categories - Treating all of the above conversion states divide by subdetector separately for leading and subleading.

To determine whether categorization is necessary or which to use as similar approach is taken to the previous sections. The significance is tested with each categorization scheme and compared with no categorization. For these tests the signal used is the $X \rightarrow \gamma\gamma$ process and with $Z \rightarrow ee$ as background. The significance is defined as $Z = N_{X\rightarrow\gamma\gamma}/\sqrt{N_{Z\rightarrow ee}}$ where $N_{X\rightarrow\gamma\gamma}$ and $N_{Z\rightarrow ee}$ are the number of events passing all selection criteria and falling within a window of $2 \times \sigma_{CB}$. For the $X \rightarrow \gamma\gamma$ signal samples, only the mass point $m_X = 90\text{GeV}$ is used. The significance defined for a single category $i$ can be written as,

$$Z_i = f_{i}^{X\rightarrow\gamma\gamma} \times N_{X\rightarrow\gamma\gamma}/\sqrt{f_{i}^{Z\rightarrow ee} \times N_{Z\rightarrow ee}}. \quad (5.4)$$

Here we have used the $f_i$ as the fraction of events in each conversion category as measured in the signal or background MC. The sum in quadrature of all conversion categories is the overall significance for that categorization scheme shown in eq. 5.5

$$Z_{\text{categories}} = \sqrt{\sum_i Z_i^2} = Z_{\text{nocategories}} \sqrt{\sum_i \left( \frac{f_{i}^{X\rightarrow\gamma\gamma}}{\sqrt{f_{i}^{Z\rightarrow ee}}} \right)^2}. \quad (5.5)$$

This allows us to use part of this equation as a figure of merit: $FoM = \sqrt{\sum_i (f_{i}^{X\rightarrow\gamma\gamma}/\sqrt{f_{i}^{Z\rightarrow ee}})^2}$. The figure of merit tells us the change in significance we expect for a particular scheme of categorization relative to having no categorization. If this number is greater than 1, the categorization is better than no categories. Tables 5.6, 5.7, and 5.8 show the results for the schemes with 4, 9 and 36 categories respectively. The FoM for each scheme is 1.71, 1.78, and 1.84 respectively. We can see from the table 5.8 that a majority of the converted photons
in both leading and subleading are of the doubleSi type. Even thought the figure of merit
for this scheme is the highest, the majority of conversion belonging to one type means we
are safe to consider fewer categories. Similarly, in the case of 4 categories, the case where
there is one converted photon, and one unconverted photon has very similar fractions for both
the leading and subleading photons being converted. Because they are so close we can merge
these categories as well. So, instead of 4 categories we would have 3: unconverted/unconverted,
(converted/unconverted + unconverted/converted), and converted/converted labeled UU, UC,
and CC respectively. It was decided ultimately to use this 3 category scheme for the analysis
as the difference in the FoM is not large and using less categories simplifies many other parts
of the analysis.

<table>
<thead>
<tr>
<th>$f^\text{str}_{Z \rightarrow ee}(%)$</th>
<th>subleading $\gamma$</th>
</tr>
</thead>
<tbody>
<tr>
<td>leading $\gamma$</td>
<td>unconverted</td>
</tr>
<tr>
<td>unconverted</td>
<td>10.2</td>
</tr>
<tr>
<td>converted</td>
<td>17.3</td>
</tr>
</tbody>
</table>

| $f^\text{str}_{H \rightarrow \gamma\gamma}(\%)$ |
|---------------------------------|--------------|
| leading $\gamma$                |
| unconverted                     | 50.5 | 21.1 |
| converted                       | 19.9 | 8.4 |

<table>
<thead>
<tr>
<th>$f_i^{X \rightarrow \gamma\gamma}/\sqrt{f^\text{str}_{Z \rightarrow ee}}$</th>
<th>subleading $\gamma$</th>
</tr>
</thead>
<tbody>
<tr>
<td>leading $\gamma$</td>
<td></td>
</tr>
<tr>
<td>unconverted</td>
<td>1.58</td>
</tr>
<tr>
<td>converted</td>
<td>0.48</td>
</tr>
</tbody>
</table>

Table 5.6: Percentage of events containing photons reconstructed as unconverted or converted
in $Z \rightarrow ee$ and $H \rightarrow \gamma\gamma$ simulated events, and $f_i^{X \rightarrow \gamma\gamma}/\sqrt{f^\text{str}_{Z \rightarrow ee}}$ per category. The Figure of Merit is 1.71.
Table 5.7: Percentage of events containing photons reconstructed as unconverted, single-track converted or double-track converted in $Z \rightarrow ee$ and $H \rightarrow \gamma\gamma$ simulated events, and $\frac{f_{X \rightarrow \gamma\gamma}}{\sqrt{f_{Z \rightarrow ee}}}$ per category. The Figure of Merit is 1.78.
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Table 5.8: Percentage of events containing photons reconstructed as unconverted or one of the five flavors of conversion in $Z \rightarrow ee$ and $H \rightarrow \gamma\gamma$ simulated events, and $f_{X\rightarrow\gamma\gamma}/\sqrt{f_{Z\rightarrow ee}}$ per category. The Figure of Merit is 1.84
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5.3.5 Selection Efficiency on Signal

The total and relative selection efficiency is summarized in Figure 5.10 where the pileup conditions corresponding to 2015+2016 and 2017 are compared. Most of the efficiency loss comes from the trigger selection (cf Table 5.4), in particular the 20 GeV or 22 GeV online cuts. The preselection cuts (i.e. asking for two loose photons with a tranverse energy higher than 22 GeV) also have a relative efficiency growing from 60 to 85% with increasing $E_T$. The offline tight efficiency is around 90%, and flat with $E_T$ as there is already a tight requirement on the trigger. The overall efficiency goes from 14%/10% for $m_X = 60$ GeV (the limit will start at 65 GeV) to 32%/24% for $m_X = 100$ GeV, for 2016 and 2017 respectively. The difference between the two years comes from the increasing pile-up which degrades the photon identification and isolation efficiencies (already included at trigger level). The total efficiency is only of 2% for $m_X = 40$ GeV, and the trigger efficiency of 4%, meaning we would need to use a different trigger with lower $E_T$ cuts in order to search below 60 GeV.

![Figure 5.10](attachment:image.png)

Figure 5.10: (a) Absolute and (b) relative selection efficiency for signal, with mc16a (2015+2016) and mc16d (2017) samples for different mass points.

The fraction of events in each category is shown in Figure 5.11. It is rather flat with the mass, with average fractions of 50%, 42% and 9% for the UU, UC and CC respectively.
Those fractions are parameterized with second order polynomial functions:

\[ f_{UU} = 5.81186.10^{-01} - 1.31921.10^{-03} * x + 4.04665.10^{-06} * x^2 \]

\[ f_{UC} = 3.51254.10^{-01} + 9.60341.10^{-04} * x - 2.99837.10^{-06} * x^2 \]

\[ f_{CC} = 1 - f_{UU} - f_{UC} \] (5.6)

Figure 5.11: Fraction of conversion categories in terms of diphoton invariant mass. A second order polynomial fit is performed. The statistical errors are binomial.

### 5.3.6 Selection Efficiency on Data

The number of selected events in data is summarized in Table 5.9. The total number of selected events is 3414135. The use of an isolation requirement in the trigger for 2017 causes fewer numbers of events to pass the preselection in 2017 compared with 2016 where no isolation was used in preselection, despite the similar integrated luminosity. After isolation is applied in the full selection the numbers are comparable.
Table 5.9: Cut flow measured in data, in the $60 < M_{\gamma\gamma} < 120$ GeV mass window, for 2015, 2016, and 2017, as well as the fraction of events in each category.

<table>
<thead>
<tr>
<th>Year</th>
<th>2015</th>
<th>2016</th>
<th>2017</th>
</tr>
</thead>
<tbody>
<tr>
<td>preselection</td>
<td>638392</td>
<td>6720477</td>
<td>4101517</td>
</tr>
<tr>
<td>tight</td>
<td>391529</td>
<td>4041009</td>
<td>2838686</td>
</tr>
<tr>
<td>isolated</td>
<td>152850</td>
<td>1564423</td>
<td>1696862</td>
</tr>
<tr>
<td>UU</td>
<td>49.9%</td>
<td>52.5%</td>
<td>47.8%</td>
</tr>
<tr>
<td>UC</td>
<td>41.0%</td>
<td>39.4%</td>
<td>42.1%</td>
</tr>
<tr>
<td>CC</td>
<td>9.1%</td>
<td>8.1%</td>
<td>9.9%</td>
</tr>
</tbody>
</table>
5.4 Signal Modeling

5.4.1 Signal Parameterization

To model the signal shape, we use a functional form that is Gaussian near the center, but has tails that are non-Gaussian. The reason for this is that energy leakage effect primarily the lower tail. This causes the distribution to be asymmetric. To model this behavior we choose a double sided crystal ball function (DSCB). This function is used several $H \rightarrow \gamma\gamma$ type analyses. Because we use the narrow width approximation (NWA) where the width of the resonance is narrower than the energy resolution of the calorimeter, this function can work for different narrow resonance models.

The double-sided Crystal Ball function is defined as

$$N \cdot \begin{cases} 
  e^{-t^2/2} & \text{if } -\alpha_{\text{Low}} \geq t \geq \alpha_{\text{High}} \\
  e^{-0.5\alpha_{\text{Low}}^2} \left( \frac{\alpha_{\text{Low}}}{\sigma_{\text{Low}}} \right)^{n_{\text{Low}}} \left( t - \alpha_{\text{Low}} \right)^{n_{\text{Low}}} & \text{if } t < -\alpha_{\text{Low}} \\
  e^{-0.5\alpha_{\text{High}}^2} \left( \frac{\alpha_{\text{High}}}{\sigma_{\text{High}}} \right)^{n_{\text{High}}} \left( t + \alpha_{\text{High}} \right)^{n_{\text{High}}} & \text{if } t > \alpha_{\text{High}},
\end{cases}$$

(5.7)

where $t = \Delta m_X / \sigma_{CB}$, $\Delta m_X = m_X - \mu_{CB}$, $N$ is a normalisation parameter, $\mu_{CB}$ is the peak of the Gaussian distribution, $\sigma_{CB}$ represents the width of the Gaussian part of the function, $\alpha_{\text{Low}}$ ($\alpha_{\text{High}}$) is the point where the Gaussian becomes a power law on the low (high) mass side, $n_{\text{Low}}$ ($n_{\text{High}}$) is the exponent of this power law. A schematic representation of the DSCB is shown in figure 5.12.

Four of the six parameters are found to be mass dependent. $n_{\text{low}}$ and $n_{\text{high}}$ are not mass dependent and so will be modeled with constants. The remaining parameters will be modeled with a first order polynomial in the mass $m_X$. To set the parameters we use the MadGraph
5. Low Mass Diphoton

ggF MC signal sample. These samples are categorized according to the 3 categories UU, UC, and CC described in section 5.3.4

To find the parameterizations as a function of $m_X$ for the DSCB we use two fits: a single point fit, and a multiple, simultaneous fit. The single point fit involves taking each mass point signal sample separately and fitting with a DSCB. The DSCB parameters for each fit are then fit with a first order polynomial, or constant in the case of $n_{low}$ and $n_{high}$ to get the parameterizations of the DSCB as a function of $m_X$. The multiple, simultaneous fit takes all of the signal mass points and fits them at once using the first order polynomials directly in the fit. The multiple fit method will better pick up the correlations between some fit parameters and so this is used as the main method to obtain the parameterization. The single mass point fits are used as a validation.

The single mass point fits for $m_X = 60, 80, \text{ and } 100$ are shown in figures 5.13, 5.14, and 5.15.
for the three conversion categories UU, UC, and CC respectively.

![Fit of the $m_{\gamma\gamma}$ distributions for $ggF$ samples at $m_X = 60$ (left), 80 (middle) and 100 GeV (right) to a double-sided Crystal Ball function for UU category. The bottom insets show the pulls in each bin (the difference between the simulated point and the fit, divided by the statistical uncertainty in each bin).](image)

![Fit of the $m_{\gamma\gamma}$ distributions for $ggF$ samples at $m_X = 60$ (left), 80 (middle) and 100 GeV (right) to a double-sided Crystal Ball function for UC category. The bottom insets show the pulls in each bin (the difference between the simulated point and the fit, divided by the statistical uncertainty in each bin).](image)

Figure 5.13: Fit of the $m_{\gamma\gamma}$ distributions for $ggF$ samples at $m_X = 60$ (left), 80 (middle) and 100 GeV (right) to a double-sided Crystal Ball function for UU category. The bottom insets show the pulls in each bin (the difference between the simulated point and the fit, divided by the statistical uncertainty in each bin).

Figure 5.14: Fit of the $m_{\gamma\gamma}$ distributions for $ggF$ samples at $m_X = 60$ (left), 80 (middle) and 100 GeV (right) to a double-sided Crystal Ball function for UC category. The bottom insets show the pulls in each bin (the difference between the simulated point and the fit, divided by the statistical uncertainty in each bin).

The results of the multiple fit are shown in table 5.10. The parameterization is shown as a function of $m_{nX} = \frac{m_X - 100}{100}$. These are the parameters that are used for the analysis. Figures 5.10-5.18 show the results of the parametrization of the multiple fit compared with the single fit points as well as the fitted line to the single fit mass points.

The correlation between all parameters in the multiple fit are shown in figure 5.19. The
strong correlation between several of the parameters shows that the multiple fit method should 
be used to make sure these relationships are fully captured by the fit.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Parameterization</th>
<th>Coefficient</th>
<th>UU</th>
<th>UC</th>
<th>CC</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta m_X$</td>
<td>$a + bm_{nX}$</td>
<td>a</td>
<td>$0.06 \pm 0.03$</td>
<td>$0.14 \pm 0.04$</td>
<td>$0.20 \pm 0.10$</td>
</tr>
<tr>
<td>$\sigma_{CB}$</td>
<td>$a + bm_{nX}$</td>
<td>a</td>
<td>$1.43 \pm 0.03$</td>
<td>$1.70 \pm 0.05$</td>
<td>$1.96 \pm 0.11$</td>
</tr>
<tr>
<td>$\alpha_{Low}$</td>
<td>$a + bm_{nX}$</td>
<td>a</td>
<td>$1.65 \pm 0.12$</td>
<td>$1.58 \pm 0.13$</td>
<td>$1.51 \pm 0.23$</td>
</tr>
<tr>
<td>$n_{Low}$</td>
<td>$a$</td>
<td>a</td>
<td>$9.85 \pm 3.75$</td>
<td>$6.74 \pm 2.34$</td>
<td>$8.04 \pm 5.91$</td>
</tr>
<tr>
<td>$\alpha_{High}$</td>
<td>$a + bm_{nX}$</td>
<td>a</td>
<td>$1.45 \pm 0.09$</td>
<td>$1.48 \pm 0.11$</td>
<td>$1.51 \pm 0.23$</td>
</tr>
<tr>
<td>$n_{High}$</td>
<td>$a$</td>
<td>a</td>
<td>$1.16 \pm 0.15$</td>
<td>$-0.05 \pm 0.16$</td>
<td>$-0.09 \pm 0.26$</td>
</tr>
</tbody>
</table>

Table 5.10: Parameterizations of the double-sided Crystal Ball function describing the signal shape, result of the multiple mass point fit at low-mass, for the three categories. $m_{nX}$ is defined as $m_{nX} = \frac{m_X - 100}{100}$. 
Figure 5.16: Result of the simultaneous fits of the $m_{\gamma\gamma}$ distributions for $ggF$ samples from 40 to 200 GeV for the $UU$ category. The four plots correspond to the parameters $\Delta m_H$, $\sigma_{CB}$, $\alpha_{low}$ and $\alpha_{high}$. The red line corresponds to the result of the multiple fit while the dashed lines correspond to the linear fit of the individual fits.
Figure 5.17: Result of the simultaneous fits of the $m_{\gamma\gamma}$ distributions for $ggF$ samples from 40 to 200 GeV for the UC category. The four plots correspond to the parameters $\Delta m_H$, $\sigma_{CB}$, $\alpha_{low}$ and $\alpha_{high}$. The red line corresponds to the result of the multiple fit while the dashed lines correspond to the linear fit of the individual fits.
Figure 5.18: Result of the simultaneous fits of the $m_{\gamma\gamma}$ distributions for $ggF$ samples from 40 to 200 GeV for the CC category. The four plots correspond to the parameters $\Delta m_H$, $\sigma_{CB}$, $\alpha_{\text{low}}$ and $\alpha_{\text{high}}$. The red line corresponds to the result of the multiple fit while the dashed lines correspond to the linear fit of the individual fits.

Figure 5.19: Correlation between different parameters of the global fit for $ggF$ samples. The index "0" indicates the constant part of the linear polynomial.
5.5 Background Modeling

One of the most important and first steps in performing any analysis is modeling the backgrounds. ATLAS sees many thousands of interactions per second but most of these are processes that we are not interested in. However, since the experimental signature can look identical to the process that we do actually care about, it is often impossible to separate the uninteresting and interesting events. This means that we will have large backgrounds in our datasets and searching for new physics often involves looking for small signals on top of the background processes.

We expect to see two main types of background, non-resonant backgrounds and resonant ones. The non-resonant backgrounds come from QCD production of photon pairs \((\gamma\gamma)\) shown in figure 5.20, photon and jet pairs \((\gamma j)\), and jet pairs \((jj)\). This background appears as a smoothly falling distribution across the entire invariant mass range and is modeled with a fit function.

![Figure 5.20: SM background processes that contribute to the non-resonant background. (a) is the \(q\bar{q}\) process and (b) is the dominant ggF process.](image)

The resonant background is due to the Drell-Yan process \(Z \rightarrow ee\) where both of the electrons are misidentified as photons shown in figure 5.21.
5.6 Non-Resonant Background

To model the non-resonant part of the background, we construct templates from background MC samples for the $\gamma\gamma$ part, and take the $\gamma$jet component from control regions in data. These two components are added together according to their respective fractions as measured in the next section.

5.6.1 Background Composition

In order to properly build the templates used to find an appropriate fit function we need to determine the relative contributions in the data from $\gamma\gamma$, $\gamma$jet, and jetjet events. This will allow us to combine the appropriate samples with the correct fraction. The $2\times2D$ sideband method is used which is described in detail in [63]. This method involves extrapolating from background control regions in the side bands of the isolation and identification variables.

To begin, a sample is prepared from the data using a modification to the Tight ID criteria where all cuts on variable measured in the first layer of the EMCal are removed except for $w_{\text{tot}}$. This working point is known as Loose'. This sample is referred to as $L'L'$. In this sample the total yield of all events, $W_{\text{tot}}^{L'L'}$, will be the sum of all $\gamma\gamma$, $\gamma$j, and j$j$ events.

$$W_{\text{tot}}^{L'L'} = W_{\gamma\gamma}^{L'L'} + W_{\gamma j}^{L'L'} + W_{j\gamma}^{L'L'} + W_{jj}^{L'L'}.$$  \hspace{1cm} (5.8)

The yields on the right of equation 5.8 are unknown. This sample is then divided into 16
orthogonal subsamples by whether each photon passes or fails the tight ID and isolation requirement. The signal region is defined by both the leading and subleading photon passing the tight ID and isolation which we denote by $TITI$. The remaining 15 control regions are where one or both photon candidates are required to fail either the ID or isolation or both. For example the region defined by the leading photon failing the tight ID and isolation is denoted $\bar{T}\bar{I}T$ and corresponds to a part of the $W_{\gamma j}^L L'$ component of equation 5.8. The total yield for any of the subsamples can be written as a function of the efficiencies to pass the tight and isolation requirement for photons, as a function of the fake rate for a jet to misidentified as a photon. There are also non-trivial correlations between the fake rates for $jj$ events and so a factor for this must also be put in. The correlation between the other components is neglected. For the $TITI$ signal region, this is,

$$N_{TITI} = W_{\gamma\gamma}^L L' \epsilon_{T1} \epsilon_{I1} \epsilon_{T2} \epsilon_{I2}$$

$$+ W_{\gamma j}^L L' \epsilon_{T1} \epsilon_{I1} f_{T1} f_{I2}$$

$$+ W_{j\gamma}^L L' f_{I1} f_{T1} \epsilon_{T2} \epsilon_{I2}$$

$$+ W_{jj}^L L' f_{I1} f_{T1} f_{I2} f_{T2} \xi_{jj}$$,

(5.9)

where

- $\epsilon_{I1}$ and $\epsilon_{I2}$ are the efficiencies of the isolation criteria of one of the six analysis under study for the leading and subleading photons respectively. They are determined from the diphoton simulation.

- $\epsilon_{T1}$ and $\epsilon_{T2}$ are the Tight identification efficiencies for the leading and subleading photons respectively, also determined from the diphoton simulation.

- $f_{I1}$ and $f_{I2}$ are the isolation fake rates for the $\gamma j$ and $j\gamma$ events, fitted directly on data.
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- \( f_{T1} \) and \( f_{T2} \) are the Tight identification fake rates for the \( \gamma j \) and \( j\gamma \) events, fitted directly on data.

- \( f'_{I1} \) and \( f'_{I2} \) are the isolation fake rates for the \( jj \) events, fitted directly on data.

- \( f'_{T1} \) and \( f'_{T2} \) are the Tight identification fake rates for the \( jj \) events, fitted directly on data.

- \( \xi_{Ijj} \) is the isolation correlation factor between the jets in the \( jj \) events, fitted directly on data.

As an example of one of the control regions, equation 5.10 shows \( TITIi \), is the subsample where the leading photon is required to pass both tight ID and isolation, but the subleading is required to fail both. The remaining 14 equations are listed in appendix A.1.

\[
N_{TITIi} = W_{\gamma\gamma}^{LL'} \epsilon_{I1} \epsilon_{T1} (1 - \epsilon_{I2})(1 - \epsilon_{T2})
+ W_{ij}^{LL'} \epsilon_{I1} \epsilon_{T1} (1 - f_{I2} - f_{T2} + f_{I2}f_{T2}) \\
+ W_{j\gamma}^{LL'} f_{I1} f_{T1} (1 - \epsilon_{I2})(1 - \epsilon_{T2}) \\
+ W_{jj}^{LL'} f'_{I1} f'_{T1} (1 - \xi_{Ijj} f'_{I2} - f'_{T2} + f'_{I2}f'_{T2} \xi_{Ijj})
\] (5.10)

With the efficiencies and isolations measured directly on the Sherpa \( \gamma\gamma \) MC sample the remaining unknowns are the fractions \( (W_{\gamma\gamma}^{LL'}, W_{\gamma j}^{LL'}, W_{j\gamma}^{LL'}, W_{jj}^{LL'}) \), the fake rates \( (f_{T1}, f_{T2}, f_{I1}, f_{I2}, f'_{T1}, f'_{T2}, f'_{I1}, f'_{I2}) \), and the \( jj \) correlation \( (\xi_{Ijj}) \). These are all determined by a fit to data. Table 5.11 shows the efficiencies measured on MC used as inputs to the fit. The results of the fits are shown in table 5.12 for each of the categories UU, UC, CC as well as the inclusive sample. The fraction of each component are shown as a function of invariant mass in figures 5.22-5.24.
Table 5.11: Isolation and identification efficiencies for true photons used as input to the 2x2D sideband method for the measurement of the diphoton purity in the [60, 120] GeV mass range of the sample passing the full selection, for the years 2015+2016 (mc16a) and 2017 (mc16d). The efficiencies are determined with respect to the leading and subleading photon candidates of true diphoton events that pass the full selection except the isolation and tight identification criteria, which are removed; the photons must pass instead the (looser) L′ identification requirements. The uncertainty arises from the MC statistics.

<table>
<thead>
<tr>
<th>Category</th>
<th>Year</th>
<th>Isolation</th>
<th>Identification</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$\epsilon_{I1}$</td>
<td>$\epsilon_{T1}$</td>
</tr>
<tr>
<td>Inclusive</td>
<td>2015+2016 (mc16a)</td>
<td>0.9291 ± 0.0004</td>
<td>0.9520 ± 0.0003</td>
</tr>
<tr>
<td></td>
<td>2017 (mc16d)</td>
<td>0.9043 ± 0.0006</td>
<td>0.9446 ± 0.0004</td>
</tr>
<tr>
<td>UU</td>
<td>2015+2016 (mc16a)</td>
<td>0.9423 ± 0.0005</td>
<td>0.9480 ± 0.0005</td>
</tr>
<tr>
<td></td>
<td>2017 (mc16d)</td>
<td>0.9174 ± 0.0008</td>
<td>0.9406 ± 0.0006</td>
</tr>
<tr>
<td>UC</td>
<td>2015+2016 (mc16a)</td>
<td>0.9198 ± 0.0006</td>
<td>0.9548 ± 0.0005</td>
</tr>
<tr>
<td></td>
<td>2017 (mc16d)</td>
<td>0.8965 ± 0.0009</td>
<td>0.9469 ± 0.0006</td>
</tr>
<tr>
<td>CC</td>
<td>2015+2016 (mc16a)</td>
<td>0.9000 ± 0.0015</td>
<td>0.9608 ± 0.0010</td>
</tr>
<tr>
<td></td>
<td>2017 (mc16d)</td>
<td>0.8741 ± 0.0021</td>
<td>0.9545 ± 0.0013</td>
</tr>
</tbody>
</table>
Table 5.12: Signal and background fraction in the mass range [60, 120] GeV, obtained with the 2x2D sideband method, for the year 2015+2016 and 2017 inclusively and in conversion category.
Figure 5.22: Data and $\gamma\gamma$, $\gamma j$, and $jj$ yields for the UU category, determined by the 2x2D sideband method as a function of the diphoton mass, for the years 2015+2016 (a) and 2017 (b). The resulting fractions are shown below and for 2015+2016 (c) and 2017 (d).
Figure 5.23: Data and $\gamma\gamma$, $\gamma j$ and $jj$ yields for the UC category, determined by the 2x2D sideband method as a function of the diphoton mass, for the years 2015+2016 (a) and 2017 (b). The resulting fractions are shown below and for 2015+2016 (c) and 2017 (d).
Figure 5.24: Data and $\gamma\gamma$, $\gamma j$ and $jj$ yields for the CC category, determined by the 2x2D sideband method as a function of the diphoton mass, for the years 2015+2016 (a) and 2017 (b). The resulting fractions are shown below and for 2015+2016 (c) and 2017 (d).
5.6.2 Background Samples and Templates

Modeling the diphoton backgrounds we find in this resonance search requires the use of Monte Carlo samples in addition to samples taken from regions of data outside of our search region. Modeling the background with data only can be tricky. Before performing our analysis we cannot know whether signal exists in our selection or not. Modeling using data directly runs the risk of absorbing the signal into what we are classifying as background. For this reason, the main SM $\gamma\gamma$ backgrounds are modeled using Monte Carlo where we can be certain that the sample contains only background processes. With about 35% of the non-resonant background coming from jets faking photons, we must also account for the $\gamma j$ and $j\gamma$ background. The $jj$ background is small compared to the others and so will be ignored. Following a strategy to section 5.6.1 data is used to model the jet components of the backgrounds.

The $\gamma\gamma$ component of the background comes from the Sherpa NLO sample listed in table 5.2. Although this is a high-statistics sample, the MC/data ratio is not as high as it has been in past iterations of this analysis. This means that there is more statistical fluctuation in the sample relative to the data than we would like. The weights used in this sample can contribute to this fluctuation as well. The four main weights used for this MC sample are shown in fig 5.25. If there are weights that have large fluctuations we would like to be able to not apply them in order to minimize the uncertainty as much as possible. The photon ID SF and MC weight both have a strong dependence on $M_{\gamma\gamma}$ so we must retain these or risk sculpting the invariant mass distribution. The other two weights however, are relatively flat in $M_{\gamma\gamma}$ and what little dependence they do have is smaller than the statistical uncertainty on the MC sample (see section 5.6.3 for the distributions). These two weights can safely be removed.

While performing the spurious signal test described in the next section, it was noticed that
the normalization between the [50,90]GeV and [90,175]GeV samples was slightly different. The spurious signal test tends to be very sensitive to these kinds of shifts. Sherpa experts confirmed that the uncertainty on the cross-section of the samples is on the order of 1%. To fix this normalization, a 5th order polynomial was fit to the distribution with different weights applied to one slice of the sample. The $\chi^2/ndf$ was compared for each correction to find the best normalization between the two slices.

- nominal cross-section: $\chi^2/ndf = 241.3/114$;
- 0.99 correction: $\chi^2/ndf = 134.596/114$;
- 0.98 correction: $\chi^2/ndf = 143.65/114$.

Based on these results we apply a correction of 0.99 to the higher mass slice. The $M_{\gamma\gamma}$ distributions before and after the correction to the cross-section is applied is shown in figure 5.26.
Figure 5.26: Invariant diphoton mass distribution computed with the Sherpa NLO sample before (a) and after (b) the correction of cross-section, fitted with a fifth order polynomial function. The bottom panels show the ratio between the histograms and the fits.

In order to get the $\gamma j$ and $j\gamma$ components of the templates, we must find a control region in data that allows us to extract jet enriched samples. In the following, only 2017 data is used to construct the templates and it is assumed that the difference in shape of the 2015 and 2016 data in the invariant mass distribution is negligible.

Three control regions are tested:

- **CR1**: the two photons pass the *tight* identification cuts, one photon passes the isolation cuts and one fails it. It uses the trigger of the analysis (2g20.tight.icalovloose, corresponding to 43.0 fb$^{-1}$). It has the advantage of being very close from the signal region and the bias from inverting the cuts is minimal. But it has a high contamination from the diphoton background: the estimated fraction of $\gamma\gamma$ from MC studies is of 25%.

- **CR2**: one photon passes the *tight* identification and the isolation cuts, while the other one passes the *loose* identification but fails both the *tight* identification and the isolation cuts. It uses the 2g20.loose trigger since the online and offline *tight* identifications are very close, corresponding to 1.54 fb$^{-1}$. It has the advantage of a minimal contamination from the diphoton background (less than 10%). But this CR is far from the signal region.
so the shape could be biased by the inversion of cuts.

- CR3: the two photons pass the isolation cuts, one passes the *tight* identification, the other one passes the *loose* identification but fails the *tight* identification. It uses data from the 2g20,loose trigger and has the advantage of being close to the signal region. But the diphoton contamination is high, around 25%.

The first control region, CR1, is very close to the signal region in data which causes it to have a very high contamination of Drell-Yan background. This would make it very difficult to accurately model the $\gamma j$ shape without first removing this background. Removing the Drell-Yan would require an entire modeling study on this control region which is not feasible. This control region is not considered further.

CR3 is the closest to the signal region as it has the closest cuts. This will be used as the base line for the templates. Because CR3 uses the 2g20,loose trigger its statistics are small compared with the high-stats Sherpa $\gamma\gamma$ sample. We are unable to use CR3 to directly sum with the MC as this would cause the fluctuations in the resulting template to be very large. Instead, a ratio of the $\gamma\gamma$ MC and CR3 are taken. This ratio is then fit with a second order polynomial for the $\mathbf{UU}$ and $\mathbf{UC}$ categories, and fit with a first order polynomial for the $\mathbf{CC}$ category. This fit can then be used to reweight the $\gamma\gamma$ MC sample in order to reproduce the $\gamma j$ shape in high-statistics. This $\gamma j$ template will then be summed with the nominal $\gamma\gamma$ MC sample according to their respective purity fractions found in section 5.6.1. Figures 5.27 and 5.28 show the invariant mass shapes for the CR2 and CR3 control regions compared with the Sherpa $\gamma\gamma$ sample as well as the ratio of the control region to MC for each category.

Several templates are also created with variations. These are to assess the impact of the choice of fit function as well as the selection of the control region. No systematic uncertainties are directly applied to the templates.
• the $\gamma\gamma$ purity is increased by 5% (conservative since the relative uncertainty on the measured purity varies between 0.6 and 2.6%).

• the $\gamma\gamma$ purity is decreased by 5% (conservative since the relative uncertainty on the measured purity varies between 0.6 and 2.6%).

• the smoothing of the reducible background is changed: the Sherpa $\gamma\gamma / \text{CR3}$ ratios fitted by second(first) order polynomial are fitted by first(second) order polynomial.

• the CR2 is used instead of the CR3 for the shape of the reducible background.
Figure 5.28: $M_{\gamma\gamma}$ shapes for the data-driven CR2 and CR3 and from the diphoton MC sample for the three categories, made with 2017 data and mc16d diphoton MC for inclusive (a), UU (b), UC (c), and CC (d). All histograms are normalized to the same area. The bottom panels correspond to the ratio to the diphoton MC shape. The pink line corresponds to a first/second order polynomial fit to the CR3/$\gamma\gamma$ MC ratio.

The templates with those different variations are showed in Figure 5.29 as well as the ratio to the nominal template. The impact of the fraction of irreducible and reducible backgrounds and the choice of the function for the smoothing of the reducible background is small (2% at most), while using a different CR can give a difference up to 5%. The Spurious Signal method will be used on the five samples in order to test the robustness of the chosen function.

A closure test is also performed on the resulting templates to validate them against the signal region in data. The 2g20 loose trigger is used which corresponds to about 10% of the
Figure 5.29: Non-resonant background templates built from the irreducible and reducible components for the inclusive case and the three categories, made with the mc16a+mc16d diphoton MC and data-driven components from 2015-2017 data. The black lines correspond to the nominal template and the other ones to variations of the templates. The bottom panels show the ratios to the nominal template.

total statistics in data. A wide binning of 10GeV in $M_{\gamma\gamma}$ is also used in order to obscure any signal that may exist in this data. Figures 5.30 and 5.31 shows the templates compared with the signal region data. The Drell-Yan peak is clearly visible in the data especially in the CC category. As these templates do not model the resonant background we only check the agreement outside of the peak region. The agreement with data is good.
Figure 5.30: Invariant mass distribution from 2015+2016 data (black points), and the non-resonant template. The grey bands correspond to the envelope of all variations.
Figure 5.31: Invariant mass distribution from 2017 data (black points), and the non-resonant template. The grey bands correspond to the envelope of all variations.
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5.6.3 Spurious Signal Tests

In order to fit a function to a distribution, we first must select a function. This is not always as easy as it seems since most times in physics we don’t know what the underlying process is generating the distribution. Without an exact functional form, we must choose one that we think approximates the distribution well. A common test in ATLAS for the suitability of a function to model the data is the spurious signal test. This involves fitting your chosen background function plus a function for your signal model on a Monte Carlo sample that contains only background events. Because we have limited resources with which to generate the MC samples, there will be statistical fluctuation in them. This will cause the signal part of the fit to report non-zero values. Because we are using a sample that has no signal in this, we call this the Spurious Signal. Obviously when we go to search for real signal in our data we would hope that our chosen functions do not give us any spurious results and so we would like them to have low values of the Spurious Signal when tested on background only samples. One way we can try to select the right function to describe the background is to test several functions against each-other, then select the one that has the lowest spurious signal, or that meets some other criteria. Which functions to test are ad-hoc and several functions that have the general behavior we see in the background distribution are chosen.

The procedure for choosing a fit function with the spurious signal test follows from the procedure used in the $H \to \gamma\gamma$ cross section analysis [62]. I will start by detailing the nominal spurious signal test that has been used in past analyses and then discuss changes made because of changes in the run 2 analysis.

- Prepare a high statistics background only sample of MC simulation.

- Fit this distribution with a signal plus background function and measure the yield from the signal part $N_{\text{spur},i}$. 
5. LOW MASS DIPOPHOTON

- For each mass point in 1 GeV steps, repeat this process.

- Find the maximum value of $N_{\text{spur},i}$ for all mass points in the range. This is called the size of the spurious signal $N_{\text{spur}}$

Normally a criteria is adopted to determine which function passes the test. In the run 1 low mass analysis, the criteria was that the size of the spurious signal $N_{\text{spur}}$ must be lower than 20\% of the background statistical uncertainty. In the standard model Higgs search, they also require that the size of $N_{\text{spur}}$ be lower than some percentage of the expected signal, however, since for this analysis there is no expected signal, this criteria was ignored.

Several fit functions may satisfy the criteria and so the one with the fewest degrees of freedom is chosen. The size of the spurious signal will also be used later in the statical model for the search as a Gaussian constraint using a width of $N_{\text{spur}}$.

The fit range for the low-mass search is $60 < m_{\gamma\gamma} < 120$ GeV. This range is selected to search below the SM Higgs mass but above the kinematic turn on caused by the 20 GeV and 22 GeV triggers used.

The MC sample used is a $\gamma\gamma$ only background sample produced with Sherpa. Several files are produced in ranges of $m_{\gamma\gamma}$ and are then summed and weighted by their cross section. This sample is then re-weighted by the shape extracted from $\gamma+$Jet data described in section 5.6.2

The fit functions chosen to test are:

- Fermi distribution: $f(x) = \exp(-x/\mu)/(1 + \exp(-(x - m_0)/\sigma))$

- Bernstein polynomials: $f(x) = \sum_{i=0}^{n} p_i u^{i}(1 - u^{i})$, where $u = (x - x_{\text{min}})/(x_{\text{max}} - x_{\text{min}})$ and $n=4$ to $n=7$

- A Landau function

- The sum of a Landau and exponential function.
For each conversion category the function is fit in the range [60, 120] GeV and the spurious signal is measured in the range [65,110] GeV. For the run 1 analysis, the function that passes the above criteria and has the fewest degrees of freedom is chosen to model the background.

For the run 2 analysis most of the spurious signal test remains unchanged, however some modifications were necessary due to a limit on the amount of Monte Carlo we were able to produce.

In the run 1 analysis because ATLAS had collected $20fb^{-1}$ of data\cite{64} compared to the $80fb^{-1}$ collected in run 2, in order to keep the spurious signal numbers near what we saw in run one, much more data would have to be generated. Limitations on the number of MC events able to be generated meant that we would have to perform this test with a much lower MC to data ratio. The standard model Higgs group uses a sample produced with a diphoton mass between 90 GeV and 175 GeV. This sample detailed above was used as the upper slice of our MC. We also needed to simulate data down to 50 GeV, so another production had to be done. Because we were the only group using this particular slice, we were not able to request the same statistics in the sample. In the end we had a MC to data ratio of $15:1$ and $50:1$ for the low and high slices respectively. A lower MC to data ratio will result in larger spurious signal numbers as the relative size of the statistical fluctuations will be higher.

The results of the spurious signal fits on the $\gamma\gamma + \gamma J$ templates discussed above are shown for the categories UU\fig{5.32}, UC\fig{5.33} and CC\fig{5.34}. The information is summarized in table\tab{5.13}, table\tab{5.14} and table\tab{5.15}. The signal function and parameterization used for these fits is the DSCB function described in section\sec{5.3}. Because the MC to data ratio is so low we are not able to find any functions that meet the nominal criteria, so we instead define a "relaxed criteria" defined by equation\eq{5.6.3}. This is used to check if the tested functions are statistically compatible with the maximum allowable
value for $S/\delta S$.

$$
\zeta_{spur} = \begin{cases} 
N_{spur} + n \star \Delta_{MC} & N_{spur} + n \star \Delta_{MC} < 0 \\
N_{spur} - n \star \Delta_{MC} & N_{spur} - n \star \Delta_{MC} > 0 \\
0 & \text{otherwise}
\end{cases}
$$

We use $n = 2$. In other words we are allowing the $2\sigma$ error bars to fall within the original range of 30% background uncertainty. Relaxing the criteria is necessary in the case of limited statistics in MC. Figure [5.39](#) shows $\zeta_{spur}$ compared with the nominal spurious signal result for each category. The relaxed criteria is included in the results tables.

Even with the case of requiring the relaxed criteria, we still do not find functions passing in each category. This inspired trying to instead perform the spurious signal test on the $M_{\gamma\gamma}$ spectrum using a relative $E_t$ (etRel) cut instead of the absolute one. The etRel is defined by dividing the photon $E_t$ by the diphoton invariant mass of the photon pair in the event. The advantage of moving to relative cuts are that the spectrum no longer shows the turn on curve from the trigger cuts which allows much easier selection of fit function. This however comes with the downside of reduced data statistics in our $M_{\gamma\gamma}$ spectrum. You can see from figure [5.35](#) the $M_{\gamma\gamma}$ spectrum for absolute $E_t$ cuts and various relative $E_t$ cuts. Selecting the cut $E_t > 22/58$ GeV completely removes the trigger turn on at the beginning of the fit range. Figure [5.36](#) show a comparison of the absolute $E_t > 22$ GeV and relative $E_t > 22/55$ GeV cuts for the full fit range in the UU category. Integrating these two spectra we get that the loss in statistics is $\sim 60\%$ going from absolute to relative cuts. Even though this seems like a large loss, the change in significance is $\sim 10\%$ (fig. [5.37](#)). We add new fit functions to the test as we are now able to consider functions that do not need to model the trigger turn on:

- Dijet: $m_{\gamma\gamma}^{-1 \log(m_{\gamma\gamma})+b2\log(m_{\gamma\gamma}^2)}(1-m_{\gamma\gamma}^{1.5})^a$
  - Dijet1: $b1 = b2 = 0$
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- Dijet2: $b^2 = 0$

- Dijet3: All coefficients included

- Exponential Poly2: $e^{(p_0 + p_1 \times m_{\gamma\gamma} + p_2 \times m_{\gamma\gamma}^2)}$

The results of the spurious signal test for the relative $E_t$ distributions are shown in figures 5.40 - 5.42 and tables 5.16 - 5.18 for each of the categories UU, UC and CC. Comparing the functions that we would select to use for the background, we can see that the relative $E_t$ cuts do perform much better as we expect (table 5.19).

The final decision on whether to use the absolute or relative cuts came down to the effect on the expected limits that could be set in each case. Figure 5.38 shows a comparison of the expected limits using the absolute $E_t$ cuts and the relative $E_t$ cuts. In the case of absolute cuts, the functions used are the ones selected and indicated in the results tables. Two different choices of functional forms are used in the relative $E_t$ limit indicated on the figure. The 5%-10% change in limit is what we would expect to see considering the similar change in significance we see. The decision was made to use the absolute $E_t$ cuts as they have slightly better limits, and don’t require changing any other parts of the analysis.

The functions chosen to model the background are the one in each category with the lowest value of $\zeta_{spur}$. This is a Landau + Exponential for the UU and UC categories, and a fifth order Bernstein polynomial for the CC.

<table>
<thead>
<tr>
<th>Category</th>
<th>function</th>
<th>$S/\delta S%$</th>
<th>$\zeta_{spur}/\delta S%$</th>
<th>$N_{spurious}$</th>
<th>Free parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>UU</td>
<td>Landau</td>
<td>-137</td>
<td>-86.1</td>
<td>-549</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Fermi 1.16e+06</td>
<td>7.46e+05</td>
<td>-1.24e+03</td>
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<td></td>
<td>LandauExp</td>
<td>-128</td>
<td>-76.7</td>
<td>-604</td>
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</tr>
<tr>
<td></td>
<td>Bern5</td>
<td>125</td>
<td>-83.4</td>
<td>701</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Bern6</td>
<td>-131</td>
<td>-88.8</td>
<td>-689</td>
<td>6</td>
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<td>Bern7</td>
<td>-142</td>
<td>-100</td>
<td>-730</td>
<td>7</td>
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</table>

Table 5.13: Spurious signal results for each of the tested functions in the UU category.
Figure 5.32: The spurious signal and the spurious signal divided by the background uncertainty with dashed lines representing 30% background uncertainty. Figures on the right include uncertainty in the UU category. The figures on the left are the same as those on the right but with no error bars shown.

<table>
<thead>
<tr>
<th>Category</th>
<th>function</th>
<th>(S/\delta S%)</th>
<th>(\zeta_{\text{spur}}/\delta S%)</th>
<th>(N_{\text{spurious}})</th>
<th>Free parameters</th>
</tr>
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<tbody>
<tr>
<td>UC</td>
<td>Landau</td>
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<td>138</td>
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<td></td>
<td>Bern5</td>
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<td>-77.2</td>
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<td>Bern6</td>
<td>-105</td>
<td>-75.2</td>
<td>-703</td>
<td>6</td>
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<tr>
<td></td>
<td>Bern7</td>
<td>-136</td>
<td>-107</td>
<td>-884</td>
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Table 5.14: Spurious signal results for each of the tested functions in the UC category.

<table>
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<tr>
<th>Category</th>
<th>function</th>
<th>(S/\delta S%)</th>
<th>(\zeta_{\text{spur}}/\delta S%)</th>
<th>(N_{\text{spurious}})</th>
<th>Free parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>CC</td>
<td>Landau</td>
<td>-121</td>
<td>-75.2</td>
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<tr>
<td></td>
<td>Fermi</td>
<td>-85</td>
<td>-37.2</td>
<td>-170</td>
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<tr>
<td></td>
<td>LandauExp</td>
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<td>-80.9</td>
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<tr>
<td></td>
<td>Bern5</td>
<td>-78.7</td>
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<tr>
<td></td>
<td>Bern6</td>
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<td>-202</td>
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<td>Bern7</td>
<td>-77.6</td>
<td>-36.3</td>
<td>-204</td>
<td>7</td>
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</table>

Table 5.15: Spurious signal results for each of the tested functions in the CC category.
Figure 5.33: The spurious signal and the spurious signal divided by the background uncertainty with dashed lines representing 30% background uncertainty. Figures on the right include uncertainty in the UC category. The figures on the left are the same as those on the right but with no error bars shown.

| Category | function | $S/\delta S\%$ | $\zeta_{spur}/\delta S\%$ | $N_{spurious}$ | Free parameters |
|----------|----------|----------------|-----------------|--------------|-----------------
| UU       | Dijet2   | -99.3          | -25.2           | 337          | 3              |
|          | Bern4    | -88.8          | -15             | -285         | 4              |
|          | Bern5    | -64            | 6.17            | 238          | 5              |
|          | ExpPoly2 | -96.5          | 48              | 326          | 2              |
|          | Landau   | -307           | -237            | -977         | 2              |
|          | Dijet1   | -531           | -482            | 1.64e+03     | 2              |
|          | Bern3    | -307           | -258            | -903         | 3              |
|          | LandauExp| 77.3           | 34.3            | 304          | 4              |
|          | Dijet3   | -114           | -30.9           | 343          | 4              |

Table 5.16: Spurious signal results for each of the tested functions in the UU category for relative $E_t$ cuts. The arrow represents the function selected to model the background in this category.
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Figure 5.34: The spurious signal and the spurious signal divided by the background uncertainty with dashed lines representing 30% background uncertainty. Figures on the right include uncertainty in the CC category. The figures on the left are the same as those on the right but with no error bars shown.

Figure 5.35: Comparison of the diphoton spectrum for absolute and relative $E_t$ cuts. The vertical line shows the beginning of the non-resonant background fit range.
Figure 5.36: Comparison of the diphoton spectrum for absolute and $E_t > 22/55\text{GeV}$ cuts for the full fit range.

Figure 5.37: Comparison of significance for absolute and relative $E_t$ cuts.

Figure 5.38: Comparison of the expected limits for absolute and relative $E_t$ cuts.
Figure 5.39: The spurious signal $S$ divided by the background uncertainty in gray, and the relaxed spurious signal $\zeta$ divided by the background uncertainty in blue as a function of $M_X$ for the UU (a), UC (b) and CC (c) categories [43].

Figure 5.40: The spurious signal and the spurious signal divided by the background uncertainty for relative $E_t$ cuts with dashed lines representing 30% background uncertainty. Figures on the right include uncertainty in the UU category. The figures on the left are the same as those on the right but with no error bars shown.
Figure 5.41: The spurious signal and the spurious signal divided by the background uncertainty for relative $E_t$ cuts with dashed lines representing 30% background uncertainty. Figures on the right include uncertainty in the UC category. The figures on the left are the same as those on the right but with no error bars shown.

<table>
<thead>
<tr>
<th>Category</th>
<th>function</th>
<th>$S/\delta S%$</th>
<th>$\zeta_{spur}/\delta S%$</th>
<th>$N_{spurious}$</th>
<th>Free parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>UC</td>
<td>ExpPoly2</td>
<td>229</td>
<td>146</td>
<td>735</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Landau</td>
<td>-215</td>
<td>164</td>
<td>-783</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Dijet1</td>
<td>-785</td>
<td>-731</td>
<td>2.47e+03</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Bern3</td>
<td>-143</td>
<td>-104</td>
<td>-407</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Dijet2</td>
<td>-171</td>
<td>-105</td>
<td>723</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Bern4</td>
<td>-99.2</td>
<td>-33.3</td>
<td>-312</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>LandauExp</td>
<td>-110</td>
<td>-44.1</td>
<td>-424</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Dijet3</td>
<td>-175</td>
<td>111</td>
<td>764</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Bern5</td>
<td>97.7</td>
<td>33.4</td>
<td>322</td>
<td>5</td>
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</table>

Table 5.17: Spurious signal results for each of the tested functions in the UC category for relative $E_t$ cuts. The arrow represents the function selected to model the background in this category.
Figure 5.42: The spurious signal and the spurious signal divided by the background uncertainty for relative $E_t$ cuts with dashed lines representing 30% background uncertainty. Figures on the right include uncertainty in the CC category. The figures on the left are the same as those on the right but with no error bars shown.

<table>
<thead>
<tr>
<th>Category</th>
<th>function</th>
<th>$S/\delta S%$</th>
<th>$\zeta_{spur}/\delta S%$</th>
<th>$N_{spurious}$</th>
<th>Free parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>CC</td>
<td>LandauExp</td>
<td>92.9</td>
<td>24.1</td>
<td>187</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Bern5</td>
<td>57.8</td>
<td>20.1</td>
<td>102</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Landau</td>
<td>127</td>
<td>83.6</td>
<td>-158</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>ExpPoly2</td>
<td>185</td>
<td>101</td>
<td>305</td>
<td>2</td>
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<tr>
<td></td>
<td>Dijet1</td>
<td>-474</td>
<td>-424</td>
<td>817</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Bern3</td>
<td>-109</td>
<td>-37.2</td>
<td>197</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Dijet2</td>
<td>-162</td>
<td>-81.4</td>
<td>270</td>
<td>3</td>
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<tr>
<td></td>
<td>Bern4</td>
<td>-101</td>
<td>44.6</td>
<td>191</td>
<td>4</td>
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<tr>
<td></td>
<td>Dijet3</td>
<td>-184</td>
<td>-105</td>
<td>282</td>
<td>4</td>
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</tbody>
</table>

Table 5.18: Spurious signal results for each of the tested functions in the CC category for relative $E_t$ cuts. The arrow represents the function selected to model the background in this category.
Table 5.19: Comparison of selected function for the absolute and relative $E_t$ cuts. The column labeled 'Abs Cut' (Rel Cut) shows the criteria for the selected function with absolute cuts (relative cuts).
5.7 Resonant Background

The resonant background comes from the Drell-Yan process, $Z \rightarrow ee$ where both electrons are misidentified as photons. As this background is a signal like bump right in the middle of our search range, it is important to have a good description of this background especially in the CC category where this effect is largest. The background shape will be modeled with a template built from $Z \rightarrow ee$ MC. This template can then be summed with the others to create a complete background description.

The Drell-Yan background is estimated using a sample of dielectrons that are reconstructed as electrons and required to pass the tight electron ID. No isolation is applied but the $E_T$ cut used in the photon selection is used. A template is constructed from this sample and is normalized to the number of electrons faking photons we expect by using the fake rates.

The distribution of electrons faking photons has shifted kinematics compared with $Z \rightarrow ee$ events reconstructed as electrons. This is because a majority of the electrons reconstructed as photons have undergone large bremsstrahlung. This shift must be modeled for the final templates. Figure 5.43 shows the invariant mass distribution of $ee$ and $e\gamma$. To do this we use a $Z \rightarrow ee$ MC sample made of only electrons. From this sample we use events where two photons are reconstructed and that pass all selection criteria for the diphoton analysis detailed in section 5.3. These reconstructed photons must also be $\Delta R$ matched to truth electrons originating from a $Z$ decay to ensure that the photon candidates are not from final state radiation. The dielectron data distribution is then transformed so that its kinematics are closer to that of the fake photons obtained from the MC sample.
Figure 5.43: Invariant mass distributions (normalized to unity) of electron pairs (black) and pairs made of a leading truth-matched electron reconstructed as an electron and sub-leading truth-matched electron reconstructed as a photon (red) or pairs made of a leading truth-matched electron reconstructed as photon and sub-leading truth-matched electron reconstructed as an electron (blue), for unconverted (a) and converted (b) photons, obtained on a $Z \to ee$ simulation sample.

5.7.1 Transformation of Dielectron Events

The transformation is done with a Smirnov, or inverse transform [65]. This method determines the transform $\phi(x)$ needed to go from a distribution $F(x)$ to $G(x)$. First we calculate the cumulative distributions (CDF) for $F(x)$ and $G(x)$ denoted $CDF_F(x)$ and $CDF_G(x)$. If the unknown transform $\phi(x)$ takes you from $F(x) \to G(x)$ then we can write

$$CDF_F(x) = CDF_G(\phi(x)).$$  \hspace{1cm} (5.11)

Taking the inverse of $CDF_F(x)$ gives the definition of the transform,

$$\phi(x) = CDF_G^{-1}(CDF_F(x)).$$  \hspace{1cm} (5.12)

So, using the CDFs for each function defines a transformation to go from one to the other. We can use this to transform our electron distributions into photon distributions in the MC samples. Figure 5.43 shows the CDFs for $ee$, $e\gamma$, and $\gamma e$ samples obtained from $Z \to ee$ MC. These are then used to determine the transforms to take an electron distribution to the photon
distribution for both leading and subleading objects. In this case the transforms work out to be simple shifts along the $M_{\gamma\gamma}$ axis between the two CDFs. So for a particular mass point, the transform is just a horizontal shift with a magnitude of the distance from the electron CDF, to the photon CDF. We write the transforms $\delta_1$ and $\delta_2$ for the leading and subleading object respectively.

\[ m_{e'e'} = m_{ee} + \delta_1 + \delta_2 = m_{e'e} + m_{ee} - m_{ee} . \]

Once these transformations are determined, the dielectron data can be shifted so the kinematics look like the resonant background in the diphoton $M_{\gamma\gamma}$ spectrum. Figure 5.47 shows
the sum result of the transformation on both the leading and subleading electrons.

Figure 5.45: Invariant mass distributions (normalised to unity) of electron with fake photon (blue) and transformed electron (red) pairs in leading (a) and subleading (b) cases for unconverted photons, obtained on a $Z \rightarrow ee$ simulation sample.

Figure 5.46: Invariant mass distributions (normalised to unity) of electron with fake photon (blue) and transformed electron (red) pairs in leading (a) and subleading (b) cases for converted photons, obtained on a $Z \rightarrow ee$ simulation sample.

5.7.2 Electron to Photon Fake Rates

After the shapes of the leading and subleading electrons have been corrected to the photon distributions, the fake rates must be calculated. To build a template with the correct
normalizations, we must know the proper number of fakes that we can expect to occur.

To measure the electron to photon fake rates, a dielectron and a electron-photon sample are used. The fake rate is measured separately for leading ($\rho_1$) and subleading ($\rho_2$) objects. These fake rates are measured in a window around the Z-peak with a width of $6 \times \sigma_{CB}$ which is 6 times the width of a double sided crystal ball fit to the data. As this is not a MC sample, the non-resonant background must be subtracted away to give a good measure of the fake rates. This is done with a signal + background fit, where the signal portion is a double sided crystal ball and a Fermi $\times$ power law for the continuum portion of the Drell-Yan. The background
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uses a Fermi × exponential function. Figure 5.48 shows the results of the fits in the four conversion categories.

Figure 5.48: Non-resonant background subtraction from the transformed $e'e'$ invariant mass distributions performed before calculation of leading and subleading fake rates in data in four conversion categories.

The electron to photon fake rates are calculated as:

$$\rho_1 = \frac{N_{\gamma e}}{N_{e'e'}},$$

$$\rho_2 = \frac{N_{e\gamma}}{N_{e'e'}}$$

(5.13)

where as before, transformed electrons are denoted $e'$. The fake rates are first calculated on a $Z \rightarrow ee$ MC sample, shown in figure 5.49. We can see that after the transformation, the fake rate is flat in the invariant mass. This means we can use a single number for the rate and do not have to parameterize it in $M_{\gamma\gamma}$.

The final fake rates as measure on the data samples are shown in figure 5.50. The red lines represent a fit to the data inside the mass window around the Z-peak.
The total number of fake diphoton events coming from the Drell-Yan process is then

\[ N_{\gamma\gamma}^{DY} = \alpha \rho_1 \rho_2 N_{ee}^{DY} \phi(m_{ee}) , \]

where \( \phi(m_{ee}) \) represents the Smirnov transformation of mass distribution from section 5.7.1 and \( \alpha \) is a correction factor introduced to take into account the reconstruction efficiency differences between single photons and pairs of photons. This correction factor is evaluated from \( Z \rightarrow ee \) MC events and is defined as:

\[ \alpha = \frac{1}{\rho_1^{MC} \rho_2^{MC}} N_{\gamma\gamma}^{MC} / N_{ee}^{MC} . \]
Figure 5.50: Invariant mass dependence of the leading (left) and subleading (right) unconverted (top) and converted (bottom) photon fake rates, as measured in data. The red lines represent the fit in the mass window.

where two transformed electrons $e'e'$ spectrum is obtained using mapping to $\gamma\gamma$ distribution directly. Because the distributions after transformation don’t depend on the $M_{\gamma\gamma}$, the fake rate in nominator also doesn’t have mass dependence in the measured correlation factor as seen in Fig. 5.51. This factor is expected to be around 1 if the photon reconstruction does not depend on the reconstruction of the other candidate. However, since the track isolation uses the information from both photon candidates in the event to determine the correct primary vertex, there is some correlation.

Resulting normalization of the transformed distributions is obtained by rescaling $e'e'$ spectra to the levels of the expected $\gamma\gamma$ background in data using the normalization factors
Figure 5.51: Invariant mass dependence of correlation factor $\alpha$ in UU (a), UC (b), CU (c) and CC (d) categories, obtained on a $Z \rightarrow ee$ simulation sample. The red lines represent the fit in the mass window.

$$f = \alpha \rho_1 \rho_2$$ shown in Fig. 5.52

5.7.3 Resonant Background Templates

The resonant background templates are constructed from the $Z \rightarrow ee$ MC samples after kinematics transformation. These are then normalized using the fraction $f = \alpha \rho_1 \rho_2$ as described in the previous section. The resulting templates are shown in figure 5.53. Note that since we are using a 3 category scheme for this analysis, the UC and CU categories are merged in the end.
Figure 5.52: Invariant mass dependence of normalisation factor $f$ in UU (a), UC (b), CU (c) and CC (d) categories, used to obtain expected number of fake photons in data. The red lines represent the fit in the mass window.
Figure 5.53: Drell-Yan invariant mass templates (red) describing the expected diphoton background component (blue) in the UU (a), UC (b), CU (c) and CC (d) categories, obtained on a $Z \rightarrow ee$ simulation sample.
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5.7.4 Standard Model Higgs Background

The standard model Higgs represents a very signal like resonant background. With its mass as $M_{\gamma\gamma} = 125$ GeV, it is outside our search window of $[65, 110]$ GeV, however the lower tail of the Higgs peak could contribute to the background. PowHeq+Pythia8 ggF Monte Carlo is used to estimate the expected contribution to the backgrounds.

The selection described in Sec. 5.3 is applied. Table 5.20 shows the number of selected SM Higgs events for each trigger configuration as used in the data and each conversion type category. The $m_{\gamma\gamma}$ distributions are shown in Fig. 5.54 (full range) and Fig. 5.55 (mass range (60, 120) GeV).

<table>
<thead>
<tr>
<th>Sample/luminosity(fb$^{-1}$)</th>
<th>trigger</th>
<th>CC</th>
<th>CU</th>
<th>UU</th>
<th>total</th>
</tr>
</thead>
<tbody>
<tr>
<td>mc16a/15.4</td>
<td>2g20_tight</td>
<td>2.31</td>
<td>6.86</td>
<td>2.74</td>
<td>11.69</td>
</tr>
<tr>
<td>mc16a/21.5</td>
<td>2g22_tight</td>
<td>2.47</td>
<td>6.64</td>
<td>2.86</td>
<td>12.19</td>
</tr>
<tr>
<td>mc16d/43.6</td>
<td>2g20_tight_icalovloose</td>
<td>6.57</td>
<td>18.57</td>
<td>9.09</td>
<td>34.23</td>
</tr>
</tbody>
</table>

Table 5.20: The number of SM Higgs with $60 < M_{\gamma\gamma} < 120$ GeV after normalizing to the corresponding luminosity. The results are shown in three categories according to the conversion types of the photons.

Adding the Standard Model Higgs component to the predicted $\gamma\gamma + \gamma jet$ background described in Sec. 5.5, $N_{Higgs}/\sqrt{N_{bkg}}$ and $N_{Higgs}/(N_{Higgs} + N_{bkg})$ are calculated, where $N_{Higgs}$ and $N_{bkg}$ are the numbers of Standard Model Higgs and $\gamma\gamma + \gamma jet$ backgrounds respectively. Only the mc16d sample is used in this check, and $Z \rightarrow ee$ background is not considered. Results are shown in Fig. 5.56 and Fig. 5.57. One can see that the Standard Model Higgs only represents a small fraction (0.35% in the highest $M_{\gamma\gamma}$ bin, without categorization in conversion types) of the total background. Therefore, the SM Higgs contribution to the backgrounds is ignored when the final background template is built.
Figure 5.54: $M_{\gamma\gamma}$ distribution of three Standard Model Higgs MC samples. (a) CC. (b) CU. (c) UU. (d) Sum of three categories.
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Figure 5.55: $M_{\gamma\gamma}$ distribution of three Standard Model Higgs MC samples with $60 < M_{\gamma\gamma} < 120$ GeV. (a) CC. (b) CU. (c) UU. (d) Sum of three categories.
Figure 5.56: Backgrounds bias due to Standard Model Higgs. (a) CC. (b) CU. (c) UU. (d) Sum of three categories.
Figure 5.57: Fraction of Standard Model Higgs in the total background. (a) CC. (b) CU. (c) UU. (d) Sum of three categories.
5.8 Fiducial Volume and Cx Factors

5.8.1 Fiducial Volume Definition

In order to quote results in a model independent way, we must correct measurements of the cross-section. This allows the comparison with other experiments as well as allowing theorists to use the results. The correction factor will correct for detector effects such as reconstruction, identification and selection efficiencies. It is defined as the number of reconstructed signal events passing all analysis requirements, \( N_{\text{selection}} \), divided by the number of signal events generated in the fiducial volume, \( N_{\text{acceptance}} \).

\[
C_X = \frac{N_{\text{selection}}}{N_{\text{acceptance}}} \tag{5.14}
\]

The correction factor is measured on MC truth values. In order for this correction factor to provide a model independent result, it must use a definition of the fiducial volume that does not depend on the final state.

In order to choose the fiducial volume, a study on MC is done. Using samples with different production modes (ggF, VBH, WH, ZH, ttH), a wide range of kinematics, as well as different final states, we can optimize the definition of the fiducial volume.

All cuts applied on reconstructed photons are applied at truth level. Both photons should be within the detector volume, \(|\eta| < 2.37\) and not in the transition region between the barrel and endcaps, \(1.37 < |\eta| < 1.52\). The standard \(E_T\) cut is also applied, \(E_T > 22\) GeV.

Photons that are produced in final states that also have final state jets (ZH, WH, ttH) will have a much higher isolation and therefore a much lower selection efficiency. We would like the correction factor to be as stable in efficiency as possible so adding an isolation cut is required. The cut is applied on the truth calorimeter isolation variable as \( etcone20 - 0.65 * E_T < 0 \). Figure 5.58 shows the calorimeter isolation values in relation to the track isolation that is used.
in the offline kinematic selection for the analysis. Figure 5.59 shows the correction factors for different production modes with and without the truth isolation cut.

Figure 5.58: Reconstructed calorimeter isolation (a) and reconstructed track isolation (b) as a function of the truth particle isolation, for mc16a MC samples for all Higgs production modes and \( m_X = 100 \text{ GeV} \). Similarly, Figures(c) and (d) show the same for mc16d MC samples.

Also supplied for the purposes of reinterpretation of the results is the acceptance factor. Figure 5.60 shows the fiducial acceptance using the cuts described above. The parameterization of the acceptance factor as a function of \( m_X \) is given as:

\[
A_X = 0.5729 - 2.069 \cdot e^{-3.13 \cdot m_X / (100 \text{ GeV})}. \tag{5.15}
\]
5.8.2 Cx Factors in Production Modes

As it is seen in Figure 5.59b, there is a 30% difference in $C_X$ factors between ggF and ttH production modes for $m_X = 60$ GeV. This difference is mainly due to the difference in the photon $E_T$ distributions. The tight ID and photon isolation both have a dependence on $E_T$. This causes differences in the ID and isolation efficiency. The photon $E_T$ distribution
is shown in Figure 5.61. Photons from the ttH production mode have larger $E_T$ compared to those from the ggF mode. Figure 5.62 presents the Tight ID and isolation efficiencies as functions of photon $E_T$. These efficiencies increase for larger $E_T$, therefore the ttH signals have higher selection efficiencies. The contribution of the isolation efficiency to the production mode differences is small compared to that of the Tight ID. This is because the fiducial volume is defined so that the production mode differences due to isolation efficiency is reduced, as it is described above. Thus, the Tight photon ID is the main cause for the difference in $C_X$ factors for ttH and ggF signals.

Figure 5.61: Truth $E_T$ distributions for photons from different signal production modes. They are shown for (a) leading photons and (b) subleading photons which pass the pre-selection. The photons from ttH signals have larger $E_T$ compared to those from ggF signals.
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Figure 5.62: Selection efficiencies of photons with respect to truth photon $E_T$. They are shown for leading photons passing the pre-selection. (a) Tight photon ID efficiency, and (b) isolation (track and calorimeter) isolation efficiency. These efficiencies increase for larger photon $E_T$.

Figures 5.64 and 5.63 show the profile plots and correlation plots of the isolation variables with respect to the truth $E_T$ of photons. These figures demonstrate how the isolation efficiencies increase with respect to photon $E_T$.

Figure 5.63: Correlation plots of topoetcone20 and truth $E_T$ for leading photons passing the pre-selection. (a) ggF (Powheg+Pythia sample) and (b) ttH signals.
Figure 5.64: Profile plots of isolation distributions with respect to the truth $E_T$, for leading photons passing the pre-selection. The markers show the average, and the bars show the standard deviation for each $E_T$ bin. (a) Calorimeter isolation, (b) topoectone20, (c) track isolation, and (d) ptone20. Note that the large bars for ttH signal is because some of its events can take large topoetcone20 values, as it is seen in Figure 5.63b.
5.9 Systematic Uncertainties

5.9.1 Uncertainties on the Cx Factors

5.9.1.1 Production Mode

The uncertainty on the correction factor is taken as the envelope between the correction factors for the production modes farthest apart. As shown in the section 5.8, the production modes with the biggest difference in the $C_X$ factor is the nominal ggF sample and the ttH sample.

The uncertainty can be parameterized as the ratio of the fits between the two samples.

\[
\text{variation up} = \frac{0.5436 + 0.06644 \times M_{\gamma\gamma}/100 + 0.05077 \times M_{\gamma\gamma}/100 \times M_{\gamma\gamma}/100}{0.08858 + 0.7405 \times M_{\gamma\gamma}/100 - 0.1989 \times M_{\gamma\gamma}/100 \times M_{\gamma\gamma}/100} - 1
\]  

(5.16)

This uncertainty decreases from 25% at 65 GeV to 2.5% at 110 GeV.

5.9.1.2 Experimental Uncertainties

There are several uncertainties we need to account for on the calculation of $C_X$ factors associated with experimental effects. These are:

- Trigger Scale Factor
- Uncertainty from pileup
- Photon identification efficiency
- Isolation efficiency
- Photon energy resolution
- Energy scale
We saw in section 5.8 that the correction factor depends on the photon ID efficiencies as well as the isolation efficiency. These two also both depend on $E_T$, so any uncertainty on the energy resolution will also need to be accounted for.

The uncertainty on $C_X$ due to the trigger scale factors is illustrated in Figure 5.65a and parameterized as a function of $m_X$ as:

$$\text{variation up} = -0.00153 + 0.03453 \times \frac{M_{\gamma\gamma}}{100} - 0.01643 \times \frac{M_{\gamma\gamma}}{100} \times \frac{M_{\gamma\gamma}}{100}$$

$$\text{variation down} = 0.001404 - 0.03403 \times \frac{M_{\gamma\gamma}}{100} + 0.01619 \times \frac{M_{\gamma\gamma}}{100} \times \frac{M_{\gamma\gamma}}{100}$$

(5.17)

This uncertainty increases from 1.4% at 65 GeV to 1.7% at 110 GeV.

The uncertainty on $C_X$ due to the pileup is illustrated in Figure 5.65b and parameterized as a function of $m_X$ as:

$$\text{variation up} = 0.1266 - 0.1812 \times \frac{M_{\gamma\gamma}}{100} + 0.07446 \times \frac{M_{\gamma\gamma}}{100} \times \frac{M_{\gamma\gamma}}{100}$$

$$\text{variation down} = -0.1475 + 0.2319 \times \frac{M_{\gamma\gamma}}{100} - 0.1041 \times \frac{M_{\gamma\gamma}}{100} \times \frac{M_{\gamma\gamma}}{100}$$

(5.18)

This uncertainty decreases from 4.1% at 65 GeV to 1.8% at 110 GeV.

Figure 5.65: Systematic uncertainties associated with to the trigger efficiency (a) and pile-up (b) as a function of $m_X$, computed with the Madgraph ggF sample.

The uncertainty on $C_X$ due to the photon identification efficiency is illustrated in Fig-
and parameterized as a function of $m_X$ as:

\[
\text{variation up} = 0.04845 - 0.05095 \ast \frac{M_{\gamma\gamma}}{100} + 0.01904 \ast \frac{M_{\gamma\gamma}}{100} \ast \frac{M_{\gamma\gamma}}{100} \\
\text{variation down} = -0.0476 + 0.04974 \ast \frac{M_{\gamma\gamma}}{100} - 0.01854 \ast \frac{M_{\gamma\gamma}}{100} \ast \frac{M_{\gamma\gamma}}{100}
\]  
(5.19)

This uncertainty decreases from 2.3% at 65 GeV to 1.5% at 110 GeV.

The uncertainty on $C_X$ due to the photon isolation efficiency is illustrated in Figure 5.66b and parameterized as a function of $m_X$ as:

\[
\text{variation up} = 0.04495 - 0.01151 \ast \frac{M_{\gamma\gamma}}{100} + 0.006713 \ast \frac{M_{\gamma\gamma}}{100} \ast \frac{M_{\gamma\gamma}}{100} \\
\text{variation down} = -0.04398 + 0.01106 \ast \frac{M_{\gamma\gamma}}{100} - 0.00645 \ast \frac{M_{\gamma\gamma}}{100} \ast \frac{M_{\gamma\gamma}}{100}
\]  
(5.20)

This uncertainty is of %, flat as a function of $m_X$.

Figure 5.66: Systematic uncertainties associated with the photon identification (a) and isolation (b) efficiencies as a function of $m_X$, computed with the Madgraph ggF sample.

The uncertainty on $C_X$ due to the photon energy resolution is illustrated in Figure 5.67a and parameterized as a function of $m_X$ as:

\[
\text{variation up} = 0.0004173 \\
\text{variation down} = -0.01147 + 0.00934 \ast \frac{M_{\gamma\gamma}}{100} - 0.00539 \ast \frac{M_{\gamma\gamma}}{100} \ast \frac{M_{\gamma\gamma}}{100}
\]  
(5.21)

This uncertainty decreases from 0.28% at 65 GeV to 0.05% at 110 GeV.
The uncertainty on $C_X$ due to the photon energy scale is illustrated in Figure 5.67a and parameterized as a function of $m_X$ as:

\[
\text{variation up} = 0.01334 - 0.01605 \times M_{\gamma\gamma}/100 + 0.004656 \times M_{\gamma\gamma}/100 \times M_{\gamma\gamma}/100
\]

(5.22)

\[
\text{variation down} = -0.01934 + 0.02594 \times M_{\gamma\gamma}/100 - 0.008859 \times M_{\gamma\gamma}/100 \times M_{\gamma\gamma}/100
\]

This uncertainty decreases from 0.49% at 65 GeV to 0.13% at 110 GeV.

Figure 5.67: Systematic uncertainties associated with to the photon energy resolution (a) and photon energy scale (b) as a function of $m_X$, computed with the Madgraph ggF sample.

5.9.2 Signal Model

5.9.2.1 Energy Scale and Resolution

In this analysis the narrow width approximation is used which means that the shape of the signal is determined almost entirely by the energy resolution and the energy scale of the detector. The uncertainty on these two will cause some uncertainty in deviation on the mean $\Delta M_X$ and the width $\sigma_{CB}$ of the double-sided crystal ball function. The uncertainty for the energy resolution and scale are estimated by varying them up and down. The parameters $\Delta M_X$ and $\sigma_{CB}$ are refit for each variation with the multiple mass point method detailed in section 5.4, keeping all other parameters fixed to their nominal values. The uncertainty is then
calculated as the ratio of the result of the varied sample to the nominal one. The uncertainties on $\Delta m_X$ vary between 0.3 and 0.5% and the ones on the width $\sigma_{CB}$ vary between 2 and 8%.

These are summarized in table 5.21 and figure 5.68.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Parameterization</th>
<th>Coefficient</th>
<th>UU</th>
<th>UC</th>
<th>CC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy scale up</td>
<td>$a + bm_{nX}$</td>
<td>a [GeV]</td>
<td>0.55</td>
<td>0.58</td>
<td>0.56</td>
</tr>
<tr>
<td>Energy scale up</td>
<td></td>
<td>b</td>
<td>0.74</td>
<td>0.64</td>
<td>0.51</td>
</tr>
<tr>
<td>Energy scale down</td>
<td>$a + bm_{nX}$</td>
<td>a [GeV]</td>
<td>-0.44</td>
<td>-0.30</td>
<td>-0.16</td>
</tr>
<tr>
<td>Energy scale down</td>
<td></td>
<td>b</td>
<td>-0.73</td>
<td>-0.54</td>
<td>-0.46</td>
</tr>
<tr>
<td>Energy resolution up</td>
<td>$a + bm_{nX}$</td>
<td>a [GeV]</td>
<td>1.53</td>
<td>1.797</td>
<td>2.0543</td>
</tr>
<tr>
<td>Energy resolution up</td>
<td></td>
<td>b</td>
<td>0.98</td>
<td>0.99292</td>
<td>1.0513</td>
</tr>
<tr>
<td>Energy resolution down</td>
<td>$a + bm_{nX}$</td>
<td>a [GeV]</td>
<td>1.35</td>
<td>1.61</td>
<td>1.87</td>
</tr>
<tr>
<td>Energy resolution down</td>
<td></td>
<td>b</td>
<td>0.61</td>
<td>0.66</td>
<td>0.74</td>
</tr>
</tbody>
</table>

Table 5.21: Parameterizations of the double-sided Crystal Ball function describing the signal shape, result of the multiple mass point fit at low-mass, for the three categories, when the photon energy scale and resolutions are varied up and down. $m_{nX}$ is defined as $m_{nX} = \frac{m_X - 100}{100}$.

Figure 5.68: Systematic uncertainties associated with to the photon energy scale (a) and photon energy resolution (b) as a function of $m_X$, computed with the Madgraph ggF sample.
5. Low Mass Diphoton

5.9.2.2 Conversion Categories

Photon conversions happen in the material of the tracking system. The uncertainty on the number of photons that migrate from one conversion category to another is assessed by using samples that have been reconstructed with a distorted detector geometry. The Egamma group produced samples with a distorted geometry containing $\gamma$+jets that are sliced in bins of photon $E_T$ from 17 GeV to 5000 GeV. Because this is not a standard diphoton sample, we cannot use the default selection and categorization. We instead use the fraction of conversions in a particular $E_T$ and $\eta$ bin, $f(E_T, \eta)$. This fraction is shown in figure 5.69.

![Fraction of converted photons in bins of $E_T$ and $\eta$ for the (a) nominal and (b) distorted geometries.](image)

Figure 5.69: fraction of converted photons in bins of $E_T$ and $\eta$ for the (a) nominal and (b) distorted geometries.

We may then use this fraction $f(E_T, \eta)$ for each of the leading and subleading photons in the signal MC samples to get the fraction of events in each category of our signal MC adjusted for the distorted geometry. This is done by looping over all events in the signal MC and summing the fraction for the leading and subleading photons for their particular value of $E_T$ and $\eta$. For example, in the CC category we have:

$$N_{CC} = \sum_{\text{allevents}} f(E_T^{\gamma_1}, \eta^{\gamma_1}) \ast f(E_T^{\gamma_2}, \eta^{\gamma_2})$$

(5.23)
And, then for the $UU$ category we have:

\[
N_{UU} = \sum_{\text{all events}} (1 - f(E_T^{\gamma_1}, \eta^{\gamma_1})) \ast (1 - f(E_T^{\gamma_2}, \eta^{\gamma_2})) \tag{5.24}
\]

The fraction of events in each category is then just the number in that category over the total number of events in the signal sample, $f_{CC} = N_{CC} / N_{total}$ and $f_{UU} = N_{UU} / N_{total}$.

The third category $UC$ is then just $f_{CU} = 1 - f_{UU} - f_{CC}$. The fraction in each category computed with the Single Photon samples and with the $X \rightarrow \gamma\gamma$ signal samples are compared in Figure 5.70a. One can see that the fractions recomputed with the fraction in the Single Photon samples do not match the numbers computed directly with the signal sample. That is most probably due to the correlation between the two photons. But assuming that this correlation is independent from the geometry one can estimate systematic uncertainties by comparing the numbers computed with the nominal and distorted geometries, as illustrated in Figure 5.70b. Those migration uncertainties are slightly mass-dependent and this dependence with modeled by third order polynomials:

- **UU** category: $f(m_X) = -15.5734 + 0.430247 \ast m_X - 0.00427721 \ast m_X^2 + 1.33562 \ast e^{-0.05} \ast m_X^3 \%$, with an average of +4%.

- **UC** category: $f(m_X) = 8.57602 - 0.254114 \ast m_X + 0.00268446 \ast m_X^2 - 8.955 \ast e^{-0.06} \ast m_X^3 \%$, with an average of +1%.

- **UC** category: $f(m_X) = 25.4582 - 0.624664 \ast m_X + 0.00560123 \ast m_X^2 - 1.53142 \ast e^{-0.05} \ast m_X^3 \%$, with an average of -2%.

The dependence of the fraction of events in each category on the production modes was checked and is illustrated in Figure 5.71. One can see in the ratios to the ggF process that the fractions agree within statistical uncertainties. But there is a small systematic effect
and the expected limit was recomputed using more conservation values of -7.5% and +6.5% for the UU and UC categories respectively, instead of -2% and +1% on average. The expected limits with the two migration uncertainties are compared in Figure 5.72. The effect of this increased uncertainty is negligible, probably because the results are dominated by the UU category. Since the $C_X$ factors are computed inclusively the difference on the migration between category have no impact on a possible re-interpretation.

Figure 5.70: (a) Fraction of events in each category computed with the signal samples (plain markers) and using the conversion fractions from Single Photon samples (open markers). (b) Ratio between the default geometry and distorted geometry.
5. Low Mass Diphoton

Figure 5.71: (a) Fraction of events in each category computed with the signal samples for different production modes. (b) Ratio of the fraction in each category to the ggF process for the UU category (c) for the UC category (d) for the CC category.

Figure 5.72: Expected limits on the fiducial production cross-section $\sigma_{fid.} BR(X \rightarrow \gamma\gamma)$ of a narrow resonance as a function of the resonance mass $m_X$ in the 65-110 GeV mass range. The black line corresponds to the default systematics uncertainties, and the green line to more conservative migration uncertainties.
## 5.9.3 Systematics summary table

Table 5.22: Summary of the main sources of systematic uncertainty on the limit on $\sigma_{fid}$.

<table>
<thead>
<tr>
<th>Source</th>
<th>Uncertainty</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Signal yield, $C_X$ factors</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Luminosity</td>
<td>±2%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td>Trigger</td>
<td>±1.4 – 1.7%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td>Photon identification</td>
<td>±1.5 – 2.3%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td>Isolation efficiency</td>
<td>±4%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td>Photon energy scale</td>
<td>±0.13 – 0.49%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td>Photon energy resolution</td>
<td>±0.053 – 0.28%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td>Pile-up</td>
<td>±1.8 – 4.1%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td>Production mode</td>
<td>±2.4 – 25%</td>
<td>$m_X$-dependent</td>
</tr>
<tr>
<td><strong>Signal modelling</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Photon energy scale</td>
<td>±0.3 – 0.5%</td>
<td>$m_X$ and category-dependent</td>
</tr>
<tr>
<td>Photon energy resolution</td>
<td>±2 – 8%</td>
<td>$m_X$ and category-dependent</td>
</tr>
<tr>
<td><strong>Migration between categories</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Material</td>
<td>−2.0 / +1.0 / +4.1%</td>
<td>category-dependent</td>
</tr>
<tr>
<td><strong>non-resonant Background</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spurious Signal</td>
<td>604 / 496 / 181 events</td>
<td>$m_X$ and category-dependent</td>
</tr>
<tr>
<td><strong>DY Background modelling</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Peak position</td>
<td>±0.1 – 0.2%</td>
<td>category-dependent</td>
</tr>
<tr>
<td>Template shape</td>
<td>±2 – 3%</td>
<td>category-dependent</td>
</tr>
<tr>
<td>Normalisation</td>
<td>±9 – 21%</td>
<td>category-dependent</td>
</tr>
</tbody>
</table>

Figure 5.73: Summary of the systematic uncertainties affecting the signal yield or the $C_X$ factors.
5.9.4 Impact of the expected limit

The impact of the different groups of systematics uncertainties (signal, DY background, Continuous background) is tested by computing the expected limit on the fiducial production cross-section, in Figure 5.74. The largest impact comes from the Spurious Signal systematics, making this analysis systematics-limited.

Figure 5.74: Expected limits on the fiducial production cross-section $\sigma_{fid} \times BR(X \rightarrow \gamma\gamma)$ of a narrow resonance as a function of the resonance mass $m_X$ in the 65-110 GeV mass range, for different hypotheses: in red no systematic uncertainties are considered, in blue only the Spurious Signal uncertainties are considered, in green all uncertainties are considered except for the one on the $C_X$ factor, in black all uncertainties (including the ones on the signal yield and the DY background) are considered.
5. Low Mass Diphoton

5.10 Results

5.10.1 Statistical Model

The data are described using an extended PDF expressed as

$$
\mathcal{L} = \prod_{c=1}^{n_c} e^{-N_{\text{total}}^c} \prod_{i=1}^{n_{\text{data}}^c} \mathcal{L}_c(M_{\gamma\gamma}(i, c))
$$

(5.25)

where $n_c = 3$ is the number of categories, $n_{\text{data}}^c$ and $N_{\text{total}}^c$ are respectively the number of data events and the sum of the fitted numbers of events in each component in category $c$, and $M_{\gamma\gamma}(i, c)$ is the $M_{\gamma\gamma}$ value for event $i$ of category $c$. The per-event term is expressed as

$$
\mathcal{L}_c(M_{\gamma\gamma}; \sigma_{\text{fid}}, m_X, N_{uu,c}, N_{uc,c}, N_{cu,c}, N_{cc,c}, N_{\text{bkg},c}, \theta) =
$$

$$
N_{\text{X,c}}(\sigma_{\text{fid}}, m_X, \theta_{\text{X,c}}, \theta_{\text{SS}}) f_X(M_{\gamma\gamma}, m_X, x_X(m_X), \theta_{x})
$$

$$
+ N_{uu,c}(\theta_{N_{uu,c}}) f_{uu,c}(M_{\gamma\gamma}, x_{uu,c}, \theta_{uu,c})
$$

$$
+ N_{uc,c}(\theta_{N_{uc,c}}) f_{uc,c}(M_{\gamma\gamma}, x_{uc,c}, \theta_{uc,c})
$$

$$
+ N_{cu,c}(\theta_{N_{cu,c}}) f_{cu,c}(M_{\gamma\gamma}, x_{cu,c}, \theta_{cu,c})
$$

$$
+ N_{cc,c}(\theta_{N_{cc,c}}) f_{cc,c}(M_{\gamma\gamma}, x_{cc,c}, \theta_{cc,c})
$$

$$
+ N_{\text{bkg, c}} f_{\text{bkg,c}}(M_{\gamma\gamma}, c_c)
$$

(5.26)

where $\sigma_{\text{fid}}$ is the fiducial production cross-section of the new resonance of mass $m_X$; $N_{uu,c}$, $N_{uc,c}$, $N_{cu,c}$, and $N_{cc,c}$ are the number of Drell-Yan background events identified respectively as unconverted-unconverted (contributing to the UU category), unconverted-converted, converted-unconverted (both contributing to the UC category) and converted-converted (contributing to the CC category). $N_{\text{bkg,c}}$ is the fitted number of background events and $c_c$ collectively refers to the background parameters used to describe its shape; finally, $\theta$ collectively designates the nuisance parameters used to describe the systematic uncertainties. The nuisance parameters are listed below:

- $\theta_{\text{lumi}}$ : uncertainty on the integrated luminosity of the data sample.
• $\theta_{\text{trig}}, \theta_{\text{ID}}, \theta_{\text{isol}}, \theta_{\text{CX}}, \theta_{\text{PU}}$: experimental uncertainties on the signal yield.

• $\theta_{\text{ERS}}, \theta_{\text{ESS}}$: experimental uncertainties on the signal yield and shape arising from energy resolution and scale.

• $\theta_{\text{mig}}$: migration uncertainty between categories.

• $\theta_{\text{bkg},uu}, \theta_{\text{bkg},cu}, \theta_{\text{bkg},cc}$: spurious signal systematics for each category.

• $\theta_{\text{mat},uu}, \theta_{\text{mat},cu}, \theta_{\text{mat},cc}$: migration uncertainty between categories.

• $\theta_{\text{stat},uu}, \theta_{\text{stat},uc}, \theta_{\text{stat},cu}, \theta_{\text{stat},cc}$: uncertainty on the normalization of each Drell-Yan component due to limited statistics in the computation of the template. Since this computation is done independently for each component, the errors are assumed to be uncorrelated. It takes also into account the systematic effect of the variation of the mass window used to compute the electron fake rate around the Z mass.

• $\theta_{\text{gen},uu}, \theta_{\text{gen},uc}, \theta_{\text{gen},cu}, \theta_{\text{gen},cc}$: uncertainty on the normalization of each Drell-Yan component due to systematic effects of using the Sherpa generator instead of Powheg. The errors are assumed to be uncorrelated.

• $\theta_{\text{BS},uu}, \theta_{\text{BS},uc}, \theta_{\text{BS},cu}, \theta_{\text{BS},cc}$: uncertainty on the parameters of the Double-sided Crystal-Ball used as model of each Drell-Yan component computed with a bootstrap method (see Section 5.7). Since this computation is done independently for each component, the errors are assumed to be uncorrelated.
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- $\theta_{\text{peak}, \text{DY}}, \theta_{\text{sigma}, \text{DY}}$: uncertainty on the peak position and width of each Drell-Yan component due to systematic effects of the material description. These effects are assumed to be correlated between Drell-Yan components, so are described as a function of a single nuisance parameter.

- $\theta_{\text{peak}, \text{DY}}, \theta_{\text{sigma}, \text{DY}}$: systematic uncertainty on the shape of the Drell-Yan components of using the Sherpa generator instead of Powheg. These effects are assumed to be correlated between Drell-Yan components.

The quantity $N_{X,c}$ represents the number of events for the new resonance, described as the product of the total number of events (proportional to $\sigma_{\text{fid}}$) and the category fractions. This allows $\sigma_{\text{sig}}$ to be directly extracted from the fit.

The number of Drell-Yan events in each category is defined as

$$N_i = N_i^0 \exp(\sigma_{\text{stat},i} \theta_{\text{stat},i}) \exp(\sigma_{\text{mat},i} \theta_{\text{mat},i,\text{DY}}) \exp(\sigma_{\text{gen},i} \theta_{\text{gen},i,\text{DY}})$$

for each component ($i = uu, uc, cu, cc$), where $N_i^0$ is the normalization of the DY template and $\sigma_{\text{stat},i}, \sigma_{\text{mat},i}$ and $\sigma_{\text{gen},i}$ are the systematic uncertainties on the template normalization due to statistical and systematic effects (material description and alternative generator). The nuisance parameters are described above.

The PDF of each Drell-Yan component is described with double-sided Crystal Ball shape, $f_i(M_{\gamma\gamma}, x_i(\theta_i))$, where the $x_i(\theta_i)$ are the parameters of the PDF and the $\theta_i$ refer to the nuisance parameters on the Drell-Yan peak position and shape. The nominal values of the PDF parameters are obtained by fitting the nominal templates obtained in Section 5.7. The shape uncertainties have three sources: the uncertainty coming from the combined effect of limited statistics in the MC and data samples and the background subtraction used to get the template (referred as the “Bootstrap” BS, uncertainty); the uncertainty coming
from the detector material description; the uncertainty on the MC generator. The material uncertainties on the shape are obtained by fitting the up/down distorted templates for each of the two shape uncertainties and linearly interpolating the PDF parameters between the "down", "nominal" and "up" values (corresponding respectively the values -1, 0, and 1 of the \(\theta_{\text{peak},\text{DY}}\) and \(\theta_{\text{sigma},\text{DY}}\) parameters). The effect of the two shape variations are assumed to be independent. The generator uncertainties on the shape are similarly described with the \(\theta_{\text{peak},\text{DY}}\) and \(\theta_{\text{sigma},\text{DY}}\) parameters.

The continuum background PDF \(f_{\text{bkg}}(M_{\gamma\gamma}, c)\) is described by the function chosen for each category in Section 5.6.3.

### 5.10.2 Partial Unblinding

It was decided during the approval process that the background model, especially the DY one, would be validated using 10% of the data, corresponding to an integrated luminosity of 8 fb\(^{-1}\), where the events are picked up randomly. A background-only fit and a p-value scan are run and shown in Figure 5.76a and the pulls of the fits are checked carefully, as shown in Figure 5.77.

A small excess with a local significance of 2.4\(\sigma\) is observed at a mass of 90 GeV. This corresponds to a slight deviation of the nuisance parameter \(\theta_{\text{norm,uc}}\) in the systematic on the normalization of the DY background, arising from the detector material description, for uc events. After several checks nothing wrong could be found. In particular most of the excess comes from the \(cu\) events, as can be seen in Figure 5.76b.
Figure 5.75: Data distributions of $M_{\gamma\gamma}$ overlaid with background-only fits, computed with 10% of the total dataset.

Figure 5.76: Observed p-value for the background-only hypothesis, $p_0$, as a function of the resonance mass $m_X$, computed with 10% of the total dataset, (a) with the combined fit (b) by fitting the categories one by one.
Figure 5.77: Pulls of the constrained nuisance parameters obtained for a signal mass hypothesis of (a) 65 GeV, (b) 90 GeV, (c) 105 GeV, computed with 10% of the total dataset.
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5.10.3 Distributions of Invariant Mass in Data

5.10.3.1 Prefit distributions

The data distribution with the full dataset is compared to the background template described in Sections 5.6.2 and 5.7 before any fits are applied in Figure 5.78. One can see that before the fit there is already a good agreement between data and the background template within uncertainties. A small excess of data can still be noticed in the DY region, but covered by the systematic uncertainties on the normalization of the DY background.

5.10.3.2 Postfit distributions

The data distribution with the full dataset is shown in Figure 5.79, as well as the bkg-only fits. The DY peak is clearly visible, and no structure is seen in the residuals. The small excess of events in the DY region of the UC category is still visible post-fit This is related to the pull distributions in Figure 5.82b discussed below.

5.10.4 Discovery p-value

The p-value scan with the full dataset is shown in Figure 5.80. No excess is seen on data.

In order to compute the fluctuation we could except in the search range, one can use the approximate formulas to compute the global significance, since the average peak fluctuation is given by finding the value of the local significance $Z_{local}$ which gives $Z_{global} = 0$:

$$p_{global} = 1 - (1 - p_{local})^{N_{trials}}$$

$$N_{trials} = 1 + \sqrt{\pi/2} \cdot N_{indep} \cdot Z_{local} N_{indep} = \frac{\text{scan range}}{\text{peak width}}$$

which gives the following formula for $Z_{global} = 0$:

$$(1 - \Phi(Z_{local}))^{1+\sqrt{\pi/2}N_{indep}Z_{local}} = 0.5$$

(5.28)
Figure 5.78: Data distributions of $M_{\gamma\gamma}$ overlaid with the background template, for (a) the UU category, (b) the UC category and (c) the CC category. The $\gamma\gamma$ contribution is taken from the Sherpa MC samples, the $\gamma j/jj$ contribution from data-driven control regions, and the $Z \rightarrow ee$ contribution from the data-driven template. The bottom panels correspond to the ratio between data and the fit, with the total uncertainty in grey and the DY normalization uncertainty in red.

The search range is 45 GeV while the width the signal is around 1.5 GeV, which gives that we expected a $2.4\sigma$ fluctuation on average.

Figures 5.81 and 5.82 show the pulls of the constrained nuisance parameters. They do not appear to have any particular structure. The pulls of the DY normalization are a bit pulled by almost $1\sigma$. 
Figure 5.79: Data distributions of $M_{\gamma\gamma}$ overlaid with background-only fits, for (a) the inclusive selection (b) the UU category, (c) the UC category and (d) the CC category. The bottom panels correspond to the difference between data and the the fit.

5.10.5 Limit on the fiducial cross-section

The expected and observed limits are shown in Figure 5.83.
Figure 5.80: Observed p-value for the background-only hypothesis, $p_0$, in (a) logarithmic and (b) linear scale, as a function of the resonance mass $m_X$.

Figure 5.81: Pulls of the constrained nuisance parameters obtained for a signal mass hypothesis of 65 GeV, for the (a) S+B fit and (b) the background-only fit.
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(a) (S+B) fit, \( m_X = 90 \) GeV

(b) B-only fit, \( m_X = 90 \) GeV

Figure 5.82: Pulls of the constrained nuisance parameters obtained for a signal mass hypothesis of 90 GeV, for the (a) S+B fit and (b) the background-only fit.

Figure 5.83: Expected and observed limits on the fiducial production cross-section \( \sigma_{\text{fid}} \cdot BR(X \to \gamma \gamma) \) of a narrow resonance as a function of the resonance mass \( m_X \) in the 65-110 GeV mass range. The green and yellow bands show the \( \pm 1\sigma \) and \( \pm 2\sigma \) uncertainties on the expected limit.
5.11 Conclusion

In this analysis we performed a search for narrow diphoton resonances in the range 65-110 GeV. No significant excesses were and a limit is set on the fiducial cross section between 80 fb and 40 fb. This analysis is limited by systematics and limited by the spurious signal systematic particularly. It is clear from figure 5.74 that the largest improvements to be made to this analysis can be made in the non-resonant background description. The spurious signal test requires large amounts of MC in order to perform well and it is no longer feasible for these samples to be produced. A new method needs to be found before the next round of analyses.
Many of the analysis and statistical methods used on ATLAS, and high energy experiments in general, are based on methods used in previous experiments where computing resources were far more limited than they are now. Although many newer methods have found their way into ATLAS, the breadth of the so called "machine learning" techniques remain relatively unexplored when compared with computer science and the data science industry. This is not completely surprising as we in the physics field tend to make much more precise measurements, and need a far greater understanding of systematic uncertainties and biases, than does the data science industry. With more complicated methods also comes more difficulty in explaining, in understanding, and in using these methods. However, the ever increasing size of the ATLAS dataset, and the requirements for better and better modeling of the data, are forcing us to increasingly turn toward these more complicated and computationally intensive methods to meet our needs.

In this chapter I will discuss Gaussian Process Regression (GPR) and an investigation into how it might be used in the low-mass diphoton analysis. Because of the very short timescale I am on to graduate, I will not be able to see GPR used in the results of the analysis, so
my intention is to document my work exploring this method. I will attempt to make this chapter as useful as possible to a student looking for an introduction to using GPR for a modeling task in their analysis. In my experience with GPR, conceptual understanding of the method has been difficult despite the quite simple mathematics of actually applying the method. Although there are many others that are much more expert in GPR than I am, I hope to be able to give a good introductory explanation that can help the reader develop a solid intuition for the mechanics of the technique. The algorithmic simplicity of GPR along with its robustness against assumptions put into a model, I think, make it a great candidate to be used for many regression problems that analysis groups face on ATLAS.

Much of this chapter is drawn from Gaussian Process for Machine Learning by Rasmussen and Williams, which is available online for free [66].

6.1 Gaussian Process Regression

There are two main viewpoints that are taken to describe the Gaussian Process, the weight-space view, and the function-space view. I find the function space view to be more intuitive and also more in line with the way we think about modeling in ATLAS, so I will concentrate on this. I am going to start with a review of some properties of multivariate Gaussian distributions and some definitions before moving into their relationship to GPR.

6.1.1 Introduction to GPR

A multivariate Gaussian (MVG) is simply a high dimension representation of a Gaussian or normal distribution. A one-dimensional Gaussian is given by equation 6.1 where $\mu$ is the mean of the distribution and $\sigma$ is the width, or standard deviation. Often, this is written as
\[ \mathcal{N}(\mu, \sigma) = \frac{1}{\sqrt{(2\pi\sigma^2)}} e^{-\frac{(x-\mu)^2}{2\sigma^2}} \] (6.1)

In the multivariate case, we just extend this to more dimensions. The \( x \) and \( \mu \) become vectors of dimension \( D \) which is also the dimension of the MVG, and \( \sigma \) becomes a matrix of size \( D \times D \) (often called the covariance matrix). We then write this as \( \mathcal{N}(x, \Sigma) \), with bold letters representing vectors and bold capitals representing matrices. MVGs have some very convenient properties, the most important of which is that the marginal distribution of a Gaussian is a Gaussian. This means that if we were to take a 2-D Gaussian and slice it along a line in the 2-D plane, the distribution along the slice would be a 1-D Gaussian. This is also true of high dimension MVGs where slices produce a Gaussian distribution of lower dimension. Figure 6.1 shows a multivariate Gaussian with \( D = 2 \) and three different marginal distributions corresponding to slicing the 2-D distribution along constant values of the \( x \) axis.

![Figure 6.1](image_url)

Figure 6.1: A 2-D Gaussian distribution with the marginalized 1-D Gaussian distributions shown projected on the left plot (a) and on their own in the right plot (b). The purple, green and yellow lines correspond to marginalization along the \( y \) axis given \( x \) values of 0, -1, -2, respectively.

Notice that depending on where we slice the 2-D distribution we get 1-D distributions with different values of \( \mu \) and \( \sigma \). We would like to be able to calculate the \( \mu \) and \( \sigma \) for the marginal
distributions given the parameters for the 2-D distribution. Luckily for Gaussians this is easy. Continuing with the setup shown in figure 6.1 we make make some more concrete definitions for the 2-D Gaussian in eq 6.2

\[
\begin{bmatrix}
  f_x \\
  f_y
\end{bmatrix} = \mathcal{N}
\begin{pmatrix}
  \begin{bmatrix}
    \mu_x \\
    \mu_y
  \end{bmatrix},
  \begin{bmatrix}
    \Sigma_{xx} & \Sigma_{xy} \\
    \Sigma_{yx} & \Sigma_{yy}
  \end{bmatrix}
\end{pmatrix}
\] (6.2)

Now we have defined this Gaussian as having dimension 2 with axes labeled \( x \) and \( y \). We would like to find the mean and width of the marginal distributions along one axis, lets say \( y \), given a value of the other axis, \( x \), written as \( \mu_{y|x} \) and \( \Sigma_{y|x} \) respectively. Eq 6.3 shows these results without proof. This is a well known result and can be found in many standard texts.

\[
\mu_{y|x} = \mu_y + \Sigma_{xy} \Sigma_{xx}^{-1} (x - \mu_x)
\]

\[
\Sigma_{y|x} = \Sigma_{yy} - \Sigma_{yx} \Sigma_{yy}^{-1} \Sigma_{yx}
\] (6.3)

\( \mu_y \) here is the \( y \)-component of the MVG mean. Often, this will be taken as 0 when used in GPR (more on this later). We will take the mean to be 0 for now (\( \mathcal{N}(\mu, \sigma) \rightarrow \mathcal{N}(0, \sigma) \))

One more point we need to make before moving on is the difference between a continuous function represented by a functional form, and a binned function that can be represented by a vector. We are used to referring to functions defined as a functional form e.g. \( f(x) = x^2 + 1 \). This is defined for all values of \( x \) in \( \mathcal{R}^1 \). When talking about binned data, it can be more convenient to describe a function as a vector \( \mathbf{f}_n = \{x_0, x_1, ..., x_n\} \). Any functional form can be described this way for a specified number of bins, but this has the advantage that any arbitrary function, including one that may not have a closed functional form, may also be represented
this way. This can be used interchangeably with the functional form when considering binned
distributions. Instead of a continuous dependent variable, the index of the vector is used in
its place.

With the tools in equation 6.3 we can begin to discuss how this can be used in GPR. As a simple example we can consider the set of data points with no uncertainty as shown
in figure 6.2. To start with GPR we need to construct a multivariate Gaussian distribution
where the number of dimensions is the number of data points in your data set (6 in the case
of figure 6.2).

![Figure 6.2: Example data points with no uncertainty.](image)

For this early example we will take the mean of this MVG, μ, to be 0. For the covariance
matrix we will need a $6 \times 6$ matrix. To fill out this matrix we will use a covariance function
$k(x, x')$, where $x$ and $x'$ are two points on the x-axis. The function $k(x, x')$ is also called the
'kernel'. With 5 data points the covariance matrix will have dimension $6 \times 6$. The role the
covariance function plays is to tell the MVG distribution how each of its axes, and in turn each
data point, are related to one another. The behavior that we would like this function to have
is that the closer two data points are together on the x-axis, the closer we would like them to
be along the y-axis. Loosely, this gives us some sense of smoothness of the data points. The
covariance function that we will use is the squared exponential shown in equation 6.4. This is
often called the Radial Basis Function (RBF) in machine learning literature. This kernel has 2 parameters, \( \sigma_f \) and \( l \). Because these are parameters of the model and not the function or distribution they are commonly called hyperparameters. These allow us to adjust the behavior of the covariance matrix. Roughly speaking, \( \sigma_f \) is the maximum allowed covariance between close points, and \( l \) is the distance at which points along the \( x \) axis become decorrelated. I will go into more detail on hyperparameters and selecting their values later. For now, let’s assume we have chosen reasonable values for \( \sigma \) and \( l \).

\[
k(x, x') = \sigma_f \exp\left( -\frac{(x - x')^2}{2l^2} \right)
\]  

(6.4)

I will denote the matrix of these \( k(x, x') \) values as \( \mathbf{K} \). We now have everything we need to construct our MVG distribution, \( \mathcal{N}(0, \mathbf{K}) \). We can now begin to make predictions at values along the \( x \) axis. This is done by marginalizing the distribution along the axis corresponding to the \( x \) value we would like to predict at (remember there is one dimension per point on the \( x \)-axis). At this point, this isn’t very interesting as the only \( x \) values we have in our matrix are the 5 points in our data. Since we are not considering any uncertainty on the points yet, this marginalization will just return the \( y \) value of the point in question (it’s easy to check this for yourself). We would like to be able to make predictions at new points on the \( x \) axis. Extending the covariance matrix to 6 dimensions will allow us to do this. The new entries in the matrix will be denoted \( \mathbf{K}_* = [k(x_*, x_1), k(x_*, x_2), ..., k(x_*, x_n)] \) where \( n \) is the number of data points and \( \mathbf{K}_{**} = k(x_*, x_*) \) with the * entries being the new \( x \) location and are added to the MVG as in equation 6.5.

\[
\begin{bmatrix}
y \\
y_*
\end{bmatrix} = \mathcal{N} \left( \begin{bmatrix}
0 \\
\mathbf{K} & \mathbf{K}_*^T \\
\mathbf{K}_* & \mathbf{K}_{**}
\end{bmatrix} \right)
\]  

(6.5)
Repeating this process for many $x$ values, we can construct a high dimensional MVG with fine enough binning along the $x$ axis to look like a smooth function. For example, if we were to use 1000 equally spaced $x$ values we will get a 1000 dimension MVG where each point in this space defines one curve in our $x-y$ plane. This is the prior distribution. Figure 6.3 shows 10 random points drawn from the prior MVG distribution.

![Figure 6.3: 10 random draws from the prior distribution constructed with the squared exponential kernel.](image)

It should now be clear that every point in this MVG space represents a function whose functional form is not explicitly specified (its represented as a vector of numbers). This also means that this MVG can be interpreted as a prior probability distribution over functions. To perform regression, what we really want is the posterior distribution. This will be a distribution over likely functions that describe the data that the MVG is conditioned on. To get the posterior distribution, we can return to equation 6.3. This is easy to extend to more than two dimensions to cover our example case. This is shown in equation 6.6 for the mean and variance of the posterior distribution along the $x_*$ axis.

$$y_* = K_* K^{-1} y$$

$$\text{var}(y_*) = K_{**} - K_* K^{-1} K_*^T$$

(6.6)

Now we are able to get the marginalized distribution along an axis of the MVG corre-
sponding to any value of the $x$ axis in our data. To say this another way, at any $x$ value in our data, the MVG defines a Gaussian distribution that the value of $y(x)$ is distributed along. Figure 6.4 shows 10 random draws from the posterior distribution that have been conditioned on the example data points using 1000 points along the $x$ axis.

Figure 6.4: 10 random draws from the posterior distribution constructed with the squared exponential kernel and conditioned on the example data points. The red dashed line is the function used to sample the data from.

Figures 6.3 and 6.4 show that the Gaussian Process can be thought of a distribution over functions whose correlations are defined by the kernel $K(x, x')$. When we then consider some data, this has the effect of down-weighting parts of the posterior distribution that are not consistent with this data. In the end we are not really interested in the distribution of functions around the data, but a prediction for the most likely value that a function describing the data will take. Plotting the mean and $2\sigma$ variance at each point for the data gets us figure 6.5.

The function that the data points are taken from is shown by the red dashed line. We can see that our prediction for the mean is quite close to this function despite only having a few data points to fit. It is worth noting a few things here. No assumptions were made on the functional form or the shape we expect in the data beyond using the squared exponential
Figure 6.5: The mean (blue line) and $2\sigma$ variance (blue shaded region) of the posterior distribution constructed with the squared exponential kernel and conditioned on the example data points. The red dashed line is the function used to sample the data from.

In this example we have also not considered any uncertainty on the data points. This is reflected in the prediction made by the GP which you can see because the mean passes exactly through the data points and the variance goes to 0 at the points as well. We are unlikely to run into a situation where there is no uncertainty on our data points. For GPs the uncertainty is easy to model. In the case of the squared exponential kernel, we simply need to add the uncertainty on our measurements to the diagonal of the covariance matrix. This will make the kernel look like equation 6.7

$$k(x, x') = \sigma_f^2 \exp \left( -\frac{(x - x')^2}{2l^2} \right) + \sigma_n^2 \delta(x, x')$$

(6.7)

where $\delta(x, x')$ is the Kronecker delta. As you can see, this will mean that the diagonal of the matrix will be $\sigma_f^2 + \sigma_n^2$. To illustrate this situation, figure 6.6 shows the same generating
function as the above plots but with some random noise put in. For this simple example, a
random shift away from the function is added as well as an error bar. The uncertainty on
the points represented by the error bars are then added into the covariance function as in
equation 6.7 More points have been added to this plot as it makes the example more clear.
These points are still taken from the same underlying distribution.

![Figure 6.6: The mean (blue line) and 2σ variance (blue shaded region) of the Gaussian Process prediction to noisy data. The points have been randomly shifted away from their generating function (red dashed line) and uncertainty has been added.](image)

The mean now is not required to pass exactly through the data points and the variance
on the prediction also reflects this by staying non-zero, even close to measured values.

### 6.1.2 Kernel Hyperparameters

Just about any kernel you choose will have some number of hyperparameters in them. Our
example of the squared exponential kernel had two, \( \sigma_f \), the amplitude, and \( l \), the length scale.
Previously we had just assumed that we had reasonable choices made for these two, but now
we will discuss the ways that we can select values for the hyperparameters.
To illustrate the effect that the length scale can have on the GP prediction, figure 6.7 shows the data from the previous example with $l$ fixed to a value that is fixed to values away from the optimal value (more on what optimal means later). Setting the length scale too short results in the prediction being allowed to fluctuate up and down a lot in between the data, where as too long gives an oversmoothed prediction.

Figure 6.7: GP prediction on the data with $l = 0.3$ (a) and $l = 6$ (b).

These examples highlight the need to select appropriate values for these hyperparameters. One way to do this is to maximize the probability $P(\theta|x, y)$ where $\theta$ are our hyperparameters. In the case of two hyperparameters we have $\theta = \{\sigma_f, l\}$, but these could be any hyperparameters associated with your chosen kernel. Now, Bayes theorem tells us that maximizing this corresponds to maximizing $P(y|x, \theta)$. Because we are making the assumption that our prior probability distributions are Gaussian distributed, this is exactly our definition of the MVG space we are using for the GP (normalized to unity to make it a probability of course). The multivariate form of a normal distribution written out is

$$
\mathcal{N}(\mu, K) = \frac{1}{\sqrt{(2\pi)^n|K|}} \exp \left( -\frac{1}{2} (y - \mu)^T K^{-1} (y - \mu) \right)
$$

(6.8)
where \( n \) is the dimension (the number of data points) and \( \mu \) is the mean of the MVG which we have been setting to 0. For convenience we can take the logarithm of this distribution, and we arrive at what is called the log marginal likelihood (LML),

\[
\log P(y|x, \theta) = -\frac{1}{2} y^T K^{-1} y - \frac{1}{2} \log |K| - \frac{n}{2} \log 2\pi
\]  (6.9)

where I have set \( \mu = 0 \). The contour plot of the log marginal likelihood for our example data set is shown in fig. 6.8. We can then use our favorite minimizer to find where this distribution is extremized, and this will give us good starting values for our hyperparameters. We can see that the first term in equation (6.9) is the only one that depends on the input data points. This term will decrease the LML with increasing length scale as the GP model becomes less and less flexible and therefore less able to describe the data. The second term is often referred to as the complexity penalty. This term only depends on the covariance kernel and will increase the LML with increasing length scale. Just as the name implies, this term penalizes the likelihood for selecting more complex models with shorter length scale. See [66] for a detailed discussion of hyperparameter selection.

The values obtained this way are a great place to start, but there are a few considerations we should make when ultimately determining what values to use. One of the big advantages of the GP is that we can use the kernel and the hyperparameters to inject knowledge of our model into the prediction. As an example, let’s consider the length scale hyperparameter. Even before we go through the process of maximizing the log marginal likelihood, we may have some understanding of what we expect the behavior of our data to be. Even though our data might fluctuate up and down quite a bit, which in some cases would cause the maximization to tend toward a shorter length scale, we may know that we only expect bumps or structure beyond some characteristic length scale. We can then set a minimum value in
the optimization of this hyperparameter to ensure that the GP is fitting with some level of smoothness that we expect. This is extremely valuable in cases where you know ahead of time some of the features about the physical process you are modeling, or perhaps the response of your measurement apparatus. Selecting the correct hyperparameters can allow a lot of control over the way your GP performs while still remaining flexible enough to let the data speak for itself.

In addition to hyperparameter selection the choice of kernel can also have a similar impact on the performance of the Gaussian Process. In cases where you would like to model specific behavior, you may need to choose a kernel that has more hyperparameters, or one whose covariance has specific characteristics. As an example, consider a RBF kernel like we have used here, but instead of a single value for the length scale \( l \), we could allow the length scale to vary along the x-axis \( l(x) \). In the case where the length scale is a linear function, you get a covariance known as the Gibbs kernel. The process of choosing or designing a kernel is often referred to as kernel engineering. Reference [66] contains a lot of useful detail and discussion.
6. Gaussian Process

on several common kernels used for various regression and classification tasks.

6.2 Gaussian Process in Low Mass Diphoton

As described in Chapter 5, there are some challenges to modeling the backgrounds in the diphoton resonance search. One major challenge is the non-resonant background shape. This is currently modeled with a function that is selected by using the spurious signal test on a high statistics sample of Monte Carlo. In order to get good results from the spurious signal test, the amount of Monte Carlo that needs to be produced must increase with the amount of data collected. Already the number of events needed has become infeasible due to the high luminosity of the LHC. In addition, the turn-on curve at low invariant mass caused by the trigger requirements make appropriate functional forms difficult to find. They are also limited to only being able to model a small portion of this shape that can be described by the relatively simple functions that can be used. As we collect more data, it is becoming more clear that we are not able to easily select a non-biased description of the background. These are areas where the Gaussian Process (GP) can improve our background description.

When we first began discussing using a GP to model the diphoton non-resonant background, we took an approach similar to what another group on ATLAS uses to smooth and interpolate distributions. The dijet+ISR group looks for resonances in the dijet invariant mass spectrum in events with large initial state radiation. This group has recently begun to use GP to smooth a distribution, known as a transfer factor that is used to estimate a number of events in a signal region of the data based on a control region [67]. This factor is defined on a 2-D plane. Points of this 2-d plane are calculated in a grid and are then fit using a 2-D GP. In their case, to avoid the GP fitting any signal that might exist in the sample, they remove points in a window around where they are currently fitting for the transfer factor. This allows
them to fit across the window and not be biased by anything in the window.

For our first attempt at using the GP in diphoton, we removed a window around the search mass from the background distribution in data, similar to the dijet+ISR group’s strategy. This allowed us to fit across the window. If a signal bump exists in data, then we don’t run the risk of absorbing that into our background description since it would be completely removed. Figure 6.9 shows an example of a diphoton background distribution with a 10 GeV window removed around 125 GeV where a signal is injected. Note, this distribution has different limits than the low-mass diphoton analysis discussed earlier. During this early stage of testing there were no low-mass samples yet available. In the bottom panel the ratio of the data points to the fit shows that the signal bump injected into the sample appears to be recovered as the GP is fit only on the sidebands of the distribution where there is no signal. Although this method recovers the injected signal, it could possibly induce biases if there are effects that could cause one side of the window to move away from the background. For example, if there were two signal bumps near each other. This could cause one side of the window to be dragged upward, and even though the window is centered on a real signal, the GP would absorb a portion of it, causing a loss in sensitivity. Having two as yet unknown signals near each other in this way is very unlikely (and the kind of problem physicists would love to have), but we decided to try to leverage some of the properties of the GP to eliminate this problem.

As mentioned in the previous section, the hyperparameters can give you quite a bit of input to how you would like you GP to behave. In particular the length scale parameter of the squared exponential covariance function in equation 6.7 will be useful here. It is useful to note at this point that the length scale has units of the x-axis of the distribution you are fitting which in this case is GeV. This hyperparameter allows for the control of the scale at which structure in the GP is fit. This means that if we set an appropriate lower limit on this parameter, we should be able to cause the GP to ignore narrow bumps. In this case we have
Figure 6.9: An example of removing a window around a search point for GP fitting. The red points are diphoton MC with DSCB signal injected at 125 GeV, the blue points are the same MC with a 10 GeV window removed centered at 125 GeV, the black points are the GP fit to the blue points. The bottom panel shows the ratio of the red data points over the GP fit.

a narrow resonance of $\sim 8$ GeV for the signal and a broad, smoothly falling background. As long as the length scale is set longer than 8 GeV, we should not need to worry about the signal being absorbed in the fit. This is an example of one large advantage of using GPs. Something as simple as bounding the length scale can encode quite a bit of relevant information about the types of signals you expect to see. The procedure that we settled on in the end was to optimize the hyperparameters using the method described in Section 6.1.2 where the input used is background-only Monte Carlo. Since there is no signal in this sample, the length scale will tend toward longer values associated with the broad, smooth background. These hyperparameters can then be used in the kernel in order to perform a GP fit directly on data.

As one of the biggest problems with the spurious signal test is Monte Carlo statistics; we would like to check that this procedure of obtaining hyperparameters from MC is robust against statistics. Figure 6.10 shows distributions of the length scale obtained from optimizing on 1000 toy data sets with different numbers of events. Figure 6.10a corresponds to 1 million
events per toy and figure 6.10b to 10 million events per toy. The vertical red line at length scale $\sim 72$ GeV corresponds to the value of the length scale obtained from the optimization on the full statistics MC sample of $\sim 100$ million events. As can be seen, with lower statistics the length scale tends to optimize to a lower value. This is due to the larger fluctuations. Even with $\sim 1\%$ of the statistics we are still able to get reasonable numbers for the length scale and we should not have to worry about signal being absorbed into the background description.

![Figure 6.10](image1.png)

Figure 6.10: The length scale distribution found through LML optimization for 1000 toys. 6.10a has 1 million events per toy, 6.10b has 10 million events per toy. The red line at $l \sim 72$ GeV is the value obtained when optimized on the full MC sample with $\sim 100$ million events.

### 6.2.1 RooGP Software Development

Every analysis on ATLAS needs a statistical framework to calculate results of measurements. There are many frameworks in place, but they are all based upon the RooFit and RooStats packages [68][69]. In order be able to use Gaussian Process in the context of searches on ATLAS, a toolkit must be built that is able to interact with these frameworks. The RooFit package uses models based on classes that represent probability distribution functions (PDFs) and provides a language for constructing, composing, and minimizing parameters of them. How this is normally done is that a PDF for the signal portion and background portions are
constructed separately and then summed into a full-model PDF. In the case of a search, one can then fit this PDF to the data and extract the normalization for the signal component of the PDF (we will call this nSig). Because with a GP the prediction needs to be completely refit for every value of the signal normalization that is taken, we need to treat these PDFs slightly differently than the normal RooFit treatment.

A large portion of the work I have contributed to implementing GPs in the low-mass search has been in the development of the RooGP tool that allows RooFit to use GPs in its fitting procedure. This tool is largely designed to be an interface between the RooFit package and the Gaussian Process Regression library in the scikit-learn python package [70] using the Python language. Because of the way that RooFit performs minimization and the fact that we need to completely refit the GP at every step, the best way to perform a signal plus background (S+B) fit is to directly incorporate the signal model into RooGP. The fit procedure that the package follows is this:

- For each step in the minimization process (tested value of nSig):
  - Construct signal function containing nSig events (a DSCB in the low-mass diphoton case)
  - Subtract this signal from the data distribution.
  - Fit the GP on this signal subtracted data.
  - Add the signal function for nSig back into the result of the GP fit (modified GP).
  - Calculate the negative log likelihood \(-\log(\mathcal{L})\) of the modified GP given the data.
- Repeat until a minimum of the \(-\log(\mathcal{L})\) is found for nSig. This corresponds to the best fit value of the number of signal events.
Figure 6.11 illustrates this procedure on a background only MC sample. For these plots only a single step of the minimization process is shown. The nSig value chosen is 2000 events which is a number far from where we would expect the result because this is a background only sample (we expect close to 0). This choice is simply to make the process easy to see. Figure 6.11c shows the result of the process. You can see that when a value for nSig that is far from 0 is chosen, the fit does not agree well with the data. The minimizer will then move to the next step in the process, moving down the gradient in $-\log(L)$. If we imagine the case where the signal portion of the fit is set to have 0 events, (its normalization is 0), then we subtract nothing in the first step, add nothing in the last step and it amounts to fitting the MC sample.

The RooGP class works through a Python class which inherits from RooFit::RooAbsPdf. The base class needs only to have overloaded a constructor responsible for setting all parameters, and a function RooAbsPdf::evaluate which calculates the value of the PDF given values of its parameters. The RooGP class operates on histograms and handles all selection of hyperparameters internally. Training and fit histograms can be supplied separately in order to tune hyperparameters on a background only sample. Currently the signal model used internally by the library only supports a double-sided Crystal Ball function (DSCB) (see section 5.4) as mainly this is being developed for low-mass diphoton. Future improvements can add support for arbitrary function or the use of template histograms.

### 6.2.2 Tests with Gaussian Process

The RooGP class also supports use of a prior mean. This can be passed to the class in the form of a ROOT::TF1 or a histogram. In all of the previous examples, the mean was taken to be 0. In general the GP is very robust against this choice. Taking the prior mean to be 0 means that the prior probability distribution has a central value corresponding to a line at 0.
Figure 6.11: Illustration of the procedure for fitting nSig with RooGP. These show one step of the minimization process with nSig = 2000. Black points are background only MC. (a) The blue line is the MC with the DSCB signal function subtracted. (b) The green line is the GP fit to the blue line. (c) The red line is the GP fit with the DSCB added back in. The red line and black points do not agree well indicating that the point nSig = 2000 is far from the minimum of \(-\log(L)\). Here \(m_{yy}\) is the diphoton invariant mass.

on the fit axes, or equivalently a MVG with mean 0. When the GP is conditioned on data, this line is pulled away from 0 to the posterior prediction. In some cases the GP will perform better if the prior distribution is closer to the data distribution that will be fit. An interesting feature of the prior mean in the GP is that with increasing length scale, the prediction of the GP approaches the prior mean. In equation 6.7 and 6.3 we can see that as \(l\) gets larger, the components of the covariance matrix get smaller and the term depending on the kernel gets smaller leaving just the prior mean. Figure 6.12 shows this effect by using a first order
polynomial as the prior mean. The three plots correspond to an length scale optimized on the data, a length scale longer than that, and one that is much longer. You can see that the prediction tends toward the prior mean. The downward bump in the fit is due to the signal portion of the fit.

Figure 6.12: The GP prediction plus a DSCB function shown in blue using a first order polynomial as the prior mean shown in red. The black points are the data being fit. (a) is the fit with the length scale optimized on the data. (b) has the length scale fixed to 250. (c) has the length scale fixed to 500. Here $m_{yy}$ is the diphoton invariant mass.

Figure 6.13 shows just how flexible the Gaussian Process is for fitting complicated shapes. This is the full-range low-mass distribution showing the complicated trigger turn-on curve. In the low-mass analysis the fit is started at 60 GeV due to the difficulty in describing the turn-on shape. As you can see, the GP can easily fit complicated shapes such as this with
almost no prior assumptions. This fit is done by taking the prior mean as 0 and allowing the hyperparameters to be optimized on the data.

A logical comparison to make when considering this property of the prior mean is to compare it to the results of the functional form used in the spurious signal test from Chapter 5.6.3. For the UU category, the functional form selected was a Landau + exponential. If we just use this function as the prior mean, the GP will effectively model the residuals of the data around this function, thereby improving the prediction. If we increase the length scale used in the covariance matrix, we should see the GP approach the Landau + exponential prior. Because with the functional form we test the spurious signal, that will be a good comparison here.

Figure 6.14 shows the spurious signal at 90 GeV as a function of length scale of a background-only sample fit with a GP using a prior mean function. The dashed line corresponds to the value of the spurious signal obtained using a Landau+Exp functional form fit. The sample used is the MC background template used for the low-mass analysis in the UU category. You can clearly see that as the length scale increases the spurious signal value approaches the
Figure 6.14: Spurious signal (SS) of the GP prediction on background only MC as a function of the length scale with a signal mass at 90 GeV. The red points correspond to a GP with a Landau+exp prior mean with the red dashed line showing the spurious signal value for the Landau+exp functional form.

result obtained with the functional form. This means that as long as our length scale is long enough to not absorb signal and not so long as to be near this asymptote, the GP outperforms a functional form fit when measured with a spurious signal.

When choosing a functional form to describe the background distribution, we know that the function does not represent the underlying function that is generating the data. The function will have some bias in it. The purpose of the spurious signal test is to try to get an estimate on the size of the bias. GPs are much more flexible than a functional form and should be much less biased. We can test the bias by generating toy distributions and evaluating the distribution of spurious signal calculated for each. 2000 toys are generated and the spurious signal calculated for each in a background MC sample with a GP fit from 50 GeV to 200 GeV. The spurious signal is extracted at 125 GeV.

When there is no signal present in the data, we know that a GP can fit the shape well. Using the RooGP class we would like to test whether or not it can accurately extract the signal
Figure 6.15: The bias of the GP is tested by fitting the GP on 2000 toys generated from background MC in the range 50-200 GeV (a). The spurious signal is extracted for each toy at 125 GeV and is shown in figure (b). (c) shows the pull distribution of the spurious signal. The bias in the spurious signal for the GP is $\sim 6\%$ of the statistical uncertainty and the width of the pull distribution is $\sim 1$. 

portion of a fit if there were signal. We can do this with signal injection tests. For this I take the background templates used to model the background in the low-mass diphoton analysis. On top of these templates I inject a signal by randomly generating events according to a DSCB distribution with parameterization described in section 5.4. I then use the RooGP class and fit the signal + background GP to it and extract the signal part of the fit. Figure 6.16 shows the results of 1000 signal toys where the number of injected signal events is subtracted from the number of extracted signal events for three different signal sizes of 500 events, 1500 events, and 3000 events. The three plots in figure 6.16 correspond to different mass points that the
Figure 6.16: Signal is injected to the $UU$ background template used in low-mass diphoton. $\Delta n_{\text{Sig}}$ here is the number of signal events extracted by the GP minus the injected number of signal events. The black, red and blue lines are the results for 500, 1500, and 3000 injected events, respectively. (a) Corresponds to a signal injected at 70 GeV, (b) at 90 GeV, and (c) at 105 GeV.

signal is injected at, 70 GeV, 90 GeV, and 105 GeV. This is to test the signal extraction across the background range. These distributions all have mean near 0 which is what we expect if the GP is able to extract close to the correct number of signal events. The RMS for these test are approximately 25 for the case with 500 events and approximately 50 for the case with 3000 events which correspond to $\sim 5\%$ and $\sim 1\%$ error respectively. This shows that the GP is not significantly biased when fitting a signal + background distribution.
6.3 Conclusions

Gaussian Process is a method for regression that has found wide use in the data science and machine learning industries. It is now beginning to see use in the ATLAS community as some simpler methods of modeling are no longer accurate enough, or computationally feasible. GP’s ability to take very broad assumptions about the data that is to be modeled while still remaining flexible enough to be unbiased make it a useful tool to have for various tasks. I have begun work on implementing a GP based package that can help to solve the problems presented by the spurious signal technique currently used on the low-mass diphoton analysis. It is now impossible for the previous methods to be used on this analysis and I believe GP presents a way to move forward.

There are still several studies and refinements to be done to fully validate and prepare the GP for use in the low-mass diphoton analysis. Investigation into different kernels could be a particularly useful study. Although the GP should be a very unbiased estimation method, the use of a different kernel, one that is possibly more motivated by the actual underlying physics or detector response, could be less likely to be biased or, at the very least, can have its motivation tied directly to the physics making its use easier to justify.

Checking the bias of the GP fitting on the background distribution in the way that I have done it is very time intensive. It requires many toy MC samples to be created and the GP fit to each. I have done this for a few mass points, but in order to fully understand the performance of the method, the bias should be tested at every mass point used in the low-mass search (65-110 GeV). It may be that toys are the best way to do this, but a faster method would be very useful in the long run.

Many analyses on ATLAS make use of statistics packages derived from RooFit and RooStats but do not directly use either of these. In the case of low-mass diphoton, a package called
HistFitter is used. This package is built in C++ and uses input datacards to define the statistical models to be used in the analysis. At the time of writing this, there was only one C++ based Gaussian Process Regression library that I could find and its functionality was quite limited. The python language, on the other hand, has several high quality libraries with GP implementations. This language mismatch represents a technical hurdle to be overcome in fully implementing the a GP in an ATLAS analysis. Also, at the time of writing this, a group of students in the Diana-HEP group (http://diana-hep.org/) have been writing a python implementation of the HistFactory statistics package called PyHF [71]. This package can be used to fully implement the statistical model used in the low-mass diphoton group.
A.1 2x2D Sideband Method

Each photon candidate is classified as belonging to a category $TI, T\bar{I}, \bar{T}I$ or $\bar{T}\bar{I}$ depending on whether it fails or passes the identification and isolation criteria. A pair of candidates therefore can belong to one out of 16 categories, labeled from $TITI$ to $\bar{T}\bar{T}\bar{T}\bar{I}$, based on the categories of both photon candidates. Neglecting the photon-photon and photon-jet correlations, the number of events in each category can be written as:

$$N_{TITI} = W_{\gamma\gamma}^{L'L'} \epsilon_{T1} \epsilon_{T2}$$

$$+ W_{\gamma j}^{L'L'} \epsilon_{T1} \epsilon_{T2}$$

$$+ W_{j\gamma}^{L'L'} \epsilon_{T1} \epsilon_{T2}$$

$$+ W_{jj}^{L'L'} \epsilon_{T1} \epsilon_{T2}$$

$$N_{T\bar{T}I} = W_{\gamma\gamma}^{L'L'} \epsilon_{T1} (1 - \epsilon_{T2})$$

$$+ W_{\gamma j}^{L'L'} \epsilon_{T1} (1 - f_{T2})$$

$$+ W_{j\gamma}^{L'L'} \epsilon_{T1} (1 - f_{T2})$$

$$+ W_{jj}^{L'L'} \epsilon_{T1} (1 - \xi_{T2} f_{T2})$$

(A.1)
\[ N_{\bar{T}ITI} = W_{\gamma\gamma}' (1 - \epsilon_{I1}) \epsilon_{T1} \epsilon_{T2} \]
\[ + \quad W_{\gamma j}' (1 - \epsilon_{I1}) \epsilon_{T1} f_{I2} f_{T2} \]
\[ + \quad W_{j\gamma}' (1 - f_{I1}) f_{T1} \epsilon_{I2} \epsilon_{T2} \]
\[ + \quad W_{jj}' (1 - f_{I1} \xi_{Ijj}) f_{T1} f_{T2} (1 - \epsilon_{T2}) \]  
\( (A.3) \)

\[ N_{\bar{T}IT\bar{I}} = W_{\gamma\gamma}' (1 - \epsilon_{I1}) (1 - \epsilon_{I2}) \epsilon_{T1} \epsilon_{T2} \]
\[ + \quad W_{\gamma j}' (1 - \epsilon_{I1}) (1 - f_{I2}) \epsilon_{T1} f_{T2} \]
\[ + \quad W_{j\gamma}' (1 - \epsilon_{I2}) (1 - f_{I1}) \epsilon_{T2} f_{T1} \]
\[ + \quad W_{jj}' (1 - f_{I1} - f_{I2} + f_{I1} f_{I2} \xi_{Ijj}) f_{T1} f_{T2} \]  
\( (A.4) \)

\[ N_{\bar{T}T\bar{I}I} = W_{\gamma\gamma}' \epsilon_{I1} \epsilon_{T1} \epsilon_{I2} (1 - \epsilon_{T2}) \]
\[ + \quad W_{\gamma j}' \epsilon_{I1} \epsilon_{T1} f_{I2} (1 - f_{T2}) \]
\[ + \quad W_{j\gamma}' f_{I1} f_{T1} \epsilon_{I2} (1 - \epsilon_{T2}) \]
\[ + \quad W_{jj}' f_{I1} f_{T1} f_{I2} (1 - f_{T2}) \xi_{Ijj} \]  
\( (A.5) \)

\[ N_{\bar{T}ITI} = W_{\gamma\gamma}' \epsilon_{I1} (1 - \epsilon_{T1}) \epsilon_{I2} \epsilon_{T2} \]
\[ + \quad W_{\gamma j}' \epsilon_{I1} (1 - \epsilon_{T1}) f_{I2} f_{T2} \]
\[ + \quad W_{j\gamma}' f_{I1} (1 - f_{T1}) \epsilon_{I2} \epsilon_{T2} \]
\[ + \quad W_{jj}' f_{I1} (1 - f_{T1}) f_{I2} f_{T2} \xi_{Ijj} \]  
\( (A.6) \)
\( N_{\overline{TT}I} = W_{\gamma'y}^{LL'} \epsilon_1 \epsilon_T (1 - \epsilon_I) (1 - \epsilon_T) \)

\[ + \ W_{\gamma'j}^{LL'} \epsilon_1 \epsilon_T (1 - f_{I2} - f_{T2} + f_{I2} f_{T2}) \]

\[ + \ W_{\gamma'j}^{LL'} f_{I1} f_{T1} (1 - \epsilon_I) (1 - \epsilon_T) \]

\[ + \ W_{jj}^{LL'} f_{I1} f_{T1} (1 - \xi_{Ijj} f_{I2} - f_{T2} f_{I2} f_{T2} \xi_{Ijj}) \] (A.7)

\( N_{\overline{TT}II} = W_{\gamma'y}^{LL'} (1 - \epsilon_I) (1 - \epsilon_T) \epsilon_{I2} \epsilon_{T2} \)

\[ + \ W_{\gamma'j}^{LL'} (1 - \epsilon_I) (1 - \epsilon_T) f_{I2} f_{T2} \]

\[ + \ W_{\gamma'j}^{LL'} (1 - f_{I1} - f_{T1} + f_{I1} f_{T1}) \epsilon_{I2} \epsilon_{T2} \]

\[ + \ W_{jj}^{LL'} (1 - f_{I1} \xi_{Ijj} - f_{T1} + f_{I1} f_{T1} \xi_{Ijj}) f_{I2} f_{T2} \] (A.8)

\( N_{\overline{TITI}} = W_{\gamma'y}^{LL'} (1 - \epsilon_I) \epsilon_T (1 - \epsilon_{I2}) (1 - \epsilon_{T2}) \)

\[ + \ W_{\gamma'j}^{LL'} \epsilon_T f_{I2} (1 - f_{T2}) \]

\[ + \ W_{\gamma'j}^{LL'} f_{T1} \epsilon_{I2} (1 - \epsilon_{T2}) \]

\[ + \ W_{jj}^{LL'} (1 - f_{I1} \xi_{Ijj}) f_{I2} f_{T2} (1 - f_{T2}) \] (A.9)

\( N_{\overline{TI}TI} = W_{\gamma'y}^{LL'} \epsilon_1 (1 - \epsilon_{T1}) (1 - \epsilon_{I2}) \epsilon_{T2} \)

\[ + \ W_{\gamma'j}^{LL'} \epsilon_{T1} (1 - f_{I2}) (1 - \epsilon_{T1}) f_{T2} \]

\[ + \ W_{\gamma'j}^{LL'} f_{I1} (1 - \epsilon_I) (1 - f_{T1}) \epsilon_{T2} \]

\[ + \ W_{jj}^{LL'} f_{I1} (1 - f_{T1}) (1 - f_{I2} \xi_{Ijj}) f_{I2} \] (A.10)
\[ N_{\bar{I}ITI} = W_{\gamma\gamma}^{LL'} \epsilon_{I1} (1 - \epsilon_{T1}) \epsilon_{I2} (1 - \epsilon_{T2}) \]
\[ + \ W_{\gamma j}^{LL'} \epsilon_{I1} (1 - \epsilon_{T1}) f_{I2} (1 - f_{T2}) \]
\[ + \ W_{j\gamma}^{LL'} f_{I1} (1 - f_{T1}) \epsilon_{I2} (1 - \epsilon_{T2}) \]
\[ + \ W_{jj}^{LL'} f_{I1} f_{I2} (1 - f'_{T1} - f'_{T2} + f'_{T1} f'_{T2}) \xi_{Ij} \]
\[ \text{(A.11)} \]

\[ N_{TI\bar{T}I} = W_{\gamma\gamma}^{LL'} (1 - \epsilon_{I1}) \epsilon_{T1} (1 - \epsilon_{I2}) (1 - \epsilon_{T2}) \]
\[ + \ W_{\gamma j}^{LL'} ((1 - \epsilon_{I1}) \epsilon_{T1} (1 - f_{I2} - f_{T2} + f_{I2} f_{T2}) \]
\[ + \ W_{j\gamma}^{LL'} (1 - f_{I1}) f_{T1} (1 - \epsilon_{T2}) (1 - \epsilon_{I2}) \]
\[ + \ W_{jj}^{LL'} f_{T1} (1 - f'_{I2} - f'_{T2} - f'_{I1} + f'_{I2} f'_{T2} + f'_{T2} f'_{I1} + f'_{I2} f'_{I1} \xi_{Ij} - f'_{I1} f'_{I2} f'_{T2} \xi_{Ij}) \]
\[ \text{(A.12)} \]

\[ N_{I\bar{T}TI} = W_{\gamma\gamma}^{LL'} (1 - \epsilon_{I1}) (1 - \epsilon_{T1}) (1 - \epsilon_{I2}) \epsilon_{T2} \]
\[ + \ W_{\gamma j}^{LL'} (1 - \epsilon_{I1}) (1 - \epsilon_{T1}) (1 - f_{I2}) f_{T2} \]
\[ + \ W_{j\gamma}^{LL'} (1 - f_{I1} - f_{T1} + f_{I1} f_{T1}) (1 - \epsilon_{I2}) \epsilon_{T2} \]
\[ + \ W_{jj}^{LL'} (1 - f'_{I1} - f'_{T1} - f'_{I2} + f'_{I1} f'_{T1} + f'_{T1} f'_{I2} + f'_{I1} f'_{I2} \xi_{Ij} - f'_{I1} f'_{T1} f'_{I2} \xi_{Ij}) f'_{T2} \]
\[ \text{(A.13)} \]

\[ N_{\bar{T}I\bar{T}I} = W_{\gamma\gamma}^{LL'} \epsilon_{I1} (1 - \epsilon_{T1}) (1 - \epsilon_{I2}) (1 - \epsilon_{T2}) \]
\[ + \ W_{\gamma j}^{LL'} \epsilon_{I1} (1 - \epsilon_{T1}) (1 - f_{I2} - f_{T2} + f_{I2} f_{T2}) \]
\[ + \ W_{j\gamma}^{LL'} f_{I1} (1 - f_{T1}) (1 - \epsilon_{I2}) (1 - \epsilon_{T2}) \]
\[ + \ W_{jj}^{LL'} f'_{I1} ((1 - f'_{T1}) (1 - f'_{I2} \xi_{Ij}) - f'_{T2} (1 - f'_{T1} (1 - f'_{I2} \xi_{Ij}))) \]
\[ \text{(A.14)} \]
In those 16 equations we have 19 unknowns, six of which are inputs or are fixed:

- $\epsilon_{T1}$ and $\epsilon_{T2}$ are the TIGHT identification efficiencies for the leading and subleading photons respectively. Their values are determined from Monte Carlo simulation of the diphoton signal.

- $\epsilon_{I1}$ and $\epsilon_{I2}$ are the isolation efficiencies for the leading and subleading photons respectively, also determined with Monte-Carlo.
The other 13 unknowns are outputs from the extended 2D sideband method, namely:

- $W_{\gamma\gamma}^{L'L'}$, $W_{\gamma\gamma}^{L'j}$, $W_{j\gamma}^{L'j}$ and $W_{jj}^{L'L'}$ are the yields for the four categories of event after the loose' preselection.

- $f_{T1}$ and $f_{T2}$ are the tight identification fake rates for the leading and subleading jets in $\gamma$-jet and jet-$\gamma$ events.

- $f'_{T1}$ and $f'_{T2}$ are the tight identification fake rates for the leading and subleading jets in jet-jet events. Those fake rates are fixed to be equal to $f_{T1}$ and $f_{T2}$ respectively, as the difference between the prime and non-prime identification fake rates are much smaller than their precision.

- $f_{I1}$ and $f_{I2}$ are the isolation fake rates for the leading and subleading jets in $\gamma$-jet and jet-$\gamma$ events.

- $f'_{I1}$ and $f'_{I2}$ are the isolation fake rates for the leading and subleading jets in jet-jet events.

- $\xi_{IJ}$ is the correlation between the isolation of the jets in the $jj$ component.
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