School of Physics and Astronomy



Searching for Gravitational Waves

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Summary of thesis

I love hitting my head against a wall, so I did that for a bit.

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 \dots as observations have accumulated, the subject has become, to my mind at least, more mysterious and more inapproachable.

 $William\ Parsons,\ Third\ Earl\ of\ Rosse$



Chapter 1

Introduction

- Usual 'start of GW astronomy' stuff
- LIGO interferometer stuff. Virgo too. Soon Kagra and LIGO India
- GRBs are good candidate sources for GWs, especially after 170817A
- Triggered vs untriggered search
- There are several different search strategies for GRB GWs. They vary in the number of assumptions that they make about the source. This thesis covers two of these searches: The modelled CBC search, which assumes an inspiral, and the unmodelled Xpipeline search, which only a maximum duration and bandwidth of the GW.
- PyGRB and X are both triggered searches
- PyGRB assumptions are restrictive enough to allow a matched-filter search
- PyGRB can be made more sensitive by assuming circularly polarised GW emission, as the beam is perpendiular to the plane of the system
- Indicate the median exclusion distance for BNS systems or something like that to show how sensitive PyGRB is
- Xpipeline is a GWB search that makes minimal assumptions
- It only uses measures of coherence based on sky location
- O2 median distances
- Improved this search using machine learning
- Describe format of thesis

Chapter 2

Gravitational Wave Astronomy in a Nutshell

2.1 General Relativity

Christoffel Symbol

$$\Gamma^{\lambda}_{\mu\nu} = \frac{1}{2} g^{\lambda\rho} [\partial_{\nu} g_{\mu\rho} + \partial_{\mu} g_{\nu\rho} - \partial_{\rho} g_{\mu\nu}]$$
 (2.1)

Riemann Curvature Tensor

$$R^{\mu}_{\lambda\alpha\beta} = \partial_{\alpha}\Gamma^{\mu}_{\lambda\beta} - \partial_{\beta}\Gamma^{\mu}_{\lambda\alpha} + \Gamma^{\mu}_{\nu\alpha}\Gamma^{\nu}_{\lambda\beta} - \Gamma^{\mu}_{\nu\beta}\Gamma^{\nu}_{\lambda\alpha} \tag{2.2}$$

Ricci Tensor

$$R_{\mu\nu} = g^{\alpha\beta} R_{\alpha\mu\beta\nu} = R^{\beta}_{\mu\beta\nu} \tag{2.3}$$

Ricci Scalar

$$R = g^{\alpha\beta} R_{\alpha\beta} = R_{\beta}^{\beta} \tag{2.4}$$

The Einstein Equations

$$G_{\mu\nu} = R_{\mu\nu} - \frac{1}{2}Rg_{\mu\nu} = \frac{8\pi G}{c^4}T_{\mu\nu}$$
 (2.5)

Alternative form of the Einstein equations

$$R_{\mu\nu} = \frac{8\pi G}{c^4} \left(T_{\mu\nu} - \frac{1}{2} T g_{\mu\nu} \right) \tag{2.6}$$

2.2 Gravitational waves

General Relativity shows that spacetime can curve and move. From this, it may seem obvious that waves can travel though spacetime. But this is not a trivial fact; Einstein himself, having introduced the concept of gravitational waves in 1916, later claimed they could not exist. In this section, we will show that gravitational

waves do exist. We will do this by considering a perturbation of the flat Minkowski metric. We will then calculate the Ricci Tensor and Ricci Scalar for the perturbed metric and, with a clever choice of Gauge transformation, see that these yield a wave solution to Einstein's Equations.

2.2.1 Linearised Gravity

Let $g_{\mu\nu}$ be the metric given by adding a small perturbation $h_{\mu\nu}$ to the Minkowski metric $\eta_{\mu\nu} = \text{diag}(-1, 1, 1, 1)$. We write this metric as

$$g_{\mu\nu} = \eta_{\mu\nu} + h_{\mu\nu}, \qquad |h_{\mu\nu}| \ll 1 \ .$$
 (2.7)

To first order in h, we calculate the Ricci tensor (2.3)

$$R_{\mu\nu} = \eta^{\alpha\beta} R_{\alpha\mu\beta\nu} = \frac{1}{2} \left(\partial_{\mu} \partial^{\alpha} h_{\alpha\nu} + \partial_{\nu} \partial_{\alpha} h_{\mu}^{\alpha} - \Box h_{\mu\nu} - \partial_{\mu} \partial_{\nu} h \right)$$
 (2.8)

and the Ricci scalar (2.4)

$$R = \eta^{\mu\nu} R_{\mu\nu} = \partial_{\mu} \partial_{\alpha} h^{\mu\alpha} - \Box h, \qquad (2.9)$$

where $\Box = \partial_{\mu}\partial^{\mu}$ is the d'Alembertian operator and $h = h^{\mu}_{\mu}$ is the trace of h. From these, we find the resulting Einstein tensor is also linear in h

$$G_{\mu\nu} = R_{\mu\nu} - \frac{1}{2} R g_{\mu\nu}. \tag{2.10}$$

2.2.2 Gauge Transformation

With the following Gauge transformation, we can simplify the expression for the linear Einstein tensor

$$x^{\mu'} = x^{\mu} + \chi^{\mu}(x), \qquad \chi \ll x \ .$$
 (2.11)

As χ is small, we have $\partial_{\mu}\chi^{\nu} \ll 1$. This gives us

$$\frac{\partial x^{\mu}}{\partial x^{\alpha'}} = \delta^{\mu}_{\alpha} - \partial_{\alpha} \chi^{\mu} + \mathcal{O}(|\partial \chi|^2) . \tag{2.12}$$

Applying these results to our metric (2.7) we find

$$g_{\alpha'\beta'} = \frac{\partial x^{\mu}}{\partial x^{\alpha'}} \frac{\partial x^{\nu}}{\partial x^{\beta'}} g_{\mu\nu} = g_{\alpha\beta} - \partial_{\alpha} \chi_{\beta} - \partial_{\beta} \chi_{\alpha} . \qquad (2.13)$$

Subtracting the Minkowski metric from each side, we find

$$h_{\alpha'\beta'} = h_{\alpha\beta} - \partial_{\alpha}\chi_{\beta} - \partial_{\beta}\chi_{\alpha} . \tag{2.14}$$

We have some freedom in choosing our χ , so we impose the harmonic gauge

condition

$$\partial_{\mu}h^{\mu}_{\nu} = \frac{1}{2}\partial_{\nu}h,\tag{2.15}$$

where $h = h_{\lambda}^{\lambda}$. We can always choose a χ such that this is true. To see this, first note that $\partial'_{\mu} = \partial_{\mu} - (\partial_{\mu} \chi^{\lambda}) \partial_{\lambda}$. From this we find

$$(\partial'_{\mu}h^{\mu}_{\nu} - \frac{1}{2}\partial'_{\nu}h') \approx (\partial_{\mu}h^{\mu}_{\nu} - \frac{1}{2}\partial_{\nu}h) - \partial^{2}\chi_{\nu} . \tag{2.16}$$

Thus, if we are given an h such that 2.15 is not true, we can choose a χ such that

$$\partial^2 \chi_{\nu} = (\partial_{\mu} h_{\nu}^{\mu} - \frac{1}{2} \partial_{\nu} h) . \qquad (2.17)$$

Using 2.15, we can simplify the Ricci tensor and scalar

$$R_{\mu\nu} = -\frac{1}{2}\Box h_{\mu\nu} \tag{2.18}$$

$$R = -\frac{1}{2}\Box h \ . \tag{2.19}$$

Using these in the linearised Einstein equations (2.10) gives us

$$\Box h_{\mu\nu} - \frac{1}{2} \eta_{\mu\nu} \partial^2 h = -\frac{16\pi G}{c^4} T_{\mu\nu} . \qquad (2.20)$$

Alternatively, we can use 2.6 to write this as

$$\Box h_{\mu\nu} = -16\pi G \left(T_{\mu\nu} - \frac{1}{2} \eta_{\mu\nu} T \right). \tag{2.21}$$

In a vacuum, the right hand side of this equation becomes zero, and we can recognise as the relativistic wave equation.

2.2.3 Physical Effects of Gravitational Waves

The plane wave solution for the vacuum wave equation is

$$h_{\mu\nu}(x) = \epsilon_{\mu\nu}e^{ik_{\alpha}x^{\alpha}} \tag{2.22}$$

where $\epsilon_{\mu\nu}$, the polarisation tensor for the gravitational wave, is symmetric and constant, and k^{α} is the 4-wave-vector given by $k^{\alpha} = (\omega, \vec{k})$. Substituting this into the vacuum wave equation, we find

$$k^2 = -\omega^2 + \vec{k} = 0 \ . \tag{2.23}$$

Hence, gravitational waves travel at the speed of light.

The polarisation tensor is not arbitrary: It must satisfy the wave equation and the harmonic gauge condition. Putting the wave solution 2.22 into the harmonic gauge condition 2.15, we find that gravitational waves are transverse

$$k^{\mu}\epsilon_{\mu\nu} = 0 \ . \tag{2.24}$$

We can impose further gauge conditions as long as the harmonic gauge (and hence 2.24) is not violated. As any transformation with $\partial^2 \chi = 0$ will satisfy 2.17, and hence harmonic gauge condition, we express χ as

$$\chi_{\nu} = X_{\nu} e^{ikx} . \tag{2.25}$$

Using 2.25 and 2.22 in the transformation equation 2.14, we find the transformation equation for the polarisation tensor

$$\epsilon'_{\mu\nu} = \epsilon_{\mu\nu} - ik_{\mu}X_{\nu} - ik_{\nu}X_{\mu} . \tag{2.26}$$

Taking the trace of this, we find

$$\epsilon_{\mu}^{\prime\mu} = \epsilon_{\mu}^{\mu} - 2ik^{\mu}X_{\mu} \ . \tag{2.27}$$

Thus, we can impose the further gauge condition that the polarisation matrix be traceless by choosing coordinates such that $\epsilon^{\mu}_{\mu} = 2ik_{\mu}X^{\mu}$, and so fixing one element of X^{μ} . We can fix the other elements of X_{μ} by setting $\epsilon_{i0} = 0$ for i = 1, 2, 3. We do this using 2.26 to find

$$\epsilon'_{i0} = \epsilon_{i0} - ik_i X_0 - ik_0 X_i . (2.28)$$

Now we see that by setting $\epsilon_{i0} = ik_iX_0 - ik_0X_i$, we have $\epsilon_{i0} = 0$. As the polarisaton tensor is symmetric, we have $\epsilon_i^0 = 0$ as well.

These conditions give us that $\epsilon_{00} = 0$ as well. Using the fact that the polarisation matrix can be assumed traceless ϵ^{μ}_{μ} , the wave solution 2.22 reduces the harmonic gauge condition 2.15 to

$$\partial_{\mu}h^{\mu}_{\nu} = 0 \ . \tag{2.29}$$

As $\epsilon_{i0} = 0$, in the $\nu = 0$ case, only the $\mu = 0$ term is non-zero. Thus we are left with

$$\partial_{\mu}h_{0}^{\mu} = \epsilon_{0}^{0}ik_{0}e^{ik_{\alpha}x^{\alpha}} = 0$$
 (2.30)

Which implies $\epsilon_0^0 = 0$. As the polarisaton tensor is symmetric, we have $\epsilon_3^3 = 0$ as well.

We can make the polarisation tensor simpler by assuming the wave is traveling in the z-direction, that is $\vec{k} = (0,0,\omega)$. The transverse condition 2.24 then forces all the z-components of the polarisation matrix to be zero, $\epsilon_i^3 = \epsilon_3^i = 0$. The wave

solution 2.22 now looks like

$$h_{\mu\nu}(x) = \begin{bmatrix} 0 & 0 & 0 & 0 \\ 0 & a & b & 0 \\ 0 & b & -a & 0 \\ 0 & 0 & 0 & 0 \end{bmatrix} e^{(i\omega(z-t))} .$$

2.3 Gravitational Waves Sources

- GW generation
- Sources
 - BBH Multiple found already
 - BNS One found and Hulse-Taylor
 - NSBH
 - GRB
 - SN
 - Unknown

2.4 Gravitational Wave Detectors

Can cite soaulson and the noises paper

- Michelson Interferometer (cite R.Weiss 1972 paper to look smart)
- Measure change in output power due to interference in the arms
- Noise sources and how they are mitigated
 - Fundamental noise
 - Environmental noise
 - Instrumental noise
 - Noise budget

2.4.1 How to find something interesting in all that noise

- Autocorrelation function
- PSD and ASD (tie back to the noise budget)
- Matched filtering (for 1 detector)
- Unmodelled search
- Network statistics (coincident and coherent)

Chapter 3

Gamma-Ray Bursts

- $\bullet\,$ History of GRB search
- What GRBs are, short long, etc. with population plot
- Summary of physics that can be done with GW+GRB astronomy, as motivation
- Chapter summary

3.1 GRB Sources

3.1.1 Compact Binary Coalescence

- BNS and NSBH
- Short GRBs
- GW170817
- Inspiral, merger, ring down
- GRB cause (accretion powered GRB etc.)

3.1.2 Core Collapse

- GRB cause
- long GRBs
- Unknown signal morphology due to lack of knowledge about cause of core collapse as well as chaotic nature of the phenomenon

3.1.3 Other Sources

- Magnetars
- Millisecond Proto-magnetars
- Others

3.2 GRB Detectors

- IPN
- Swift
- Fermi

3.3 GRB Gravitational Wave Searches

- Triggered vs un-triggered
- Quick description of EM followup for un-triggered search, like BNS (not the focus of this thesis)
- Focus on Targeted search Modelled and un-modelled
- Known direction
- EM + GW allows for greater sensitivity

3.3.1 Modelled search for CBCs

- GW Waveform models (less accurate at merger)
- Matched filter search. PyGRB later

3.3.2 Unmodelled Search

• Coherent measurements. Discussed in chapter on X and MVA

3.4 GRB Astrophysics with Gravitational Waves

- Progenitors
- populations
- jet structure
- Opening angle

- Degenerate with distance
- Probably need a whole population of BNS/NSBH events to determine by comparing EM short GRBs with
- NS EoS
- Multi-messenger astronomy, Neurtinos, EM followup, BNS

Chapter 4

The Compact Binary Coalescence Search

4.1 Introduction

- CBC search for GRBs
- For short GRBs
- Integrating into PyCBC
- Coherent
- Targeted Better sensitivity but slower search method
- Matched filter
- Detected BNS with FAP of XX
- Used in O1 and O2 no other detections
- Chapter summary

4.2 PyCBC overview

- Just the parts of the CBC search that are relevant for PyGRB
- Highlight some of the differences
- All sky all time
- Searches for a bigger population i.e. not just BNS and low mass NSBH

4.2.1 Matched Filter Search

Contents of this section depends on what goes into the background chapter. I
think the mathematical formalism of matched filtering should go earlier in the
thesis

4.2.2 Consistency Checks

Chisq tests

- Frequency bins
- Template bank
- Autocorrelation
- Need to learn about the effective degrees of freedom stuff.
- Does the frequency bin chi2 test use other waveforms? How?

4.2.3 PyCBC Results

- Detections made in O1 and O2 Use catalog paper
- Sensitivity of search Useful later for comparison between PyCBC and PyGRB

4.3 PyGRB

The PyCBC pipeline discussed above is an all-time, all-sky search. Some CBC signals (those with one neutron star component and either a low mass black hole or another neutron star as the second component) are expected to produce a short GRB (see chapter 3). Targeted searches, where the time and sky position of the source is known, are not only computationally cheaper, but can make use of improved methods of GW detection that are too computationally expensive to use on an all-sky, all-time search.

One of these improved methods, used by PyGRB, is to calculate the *Coherent SNR*. Knowing the sky position and time a candidate GW source, as well as the antenna response function and PSD of each detector in the network allows us to calculate the amount of signal power expected in each detector. For example, detectors are most sensitive to GW sources that are directly overhead. This means a GW source that is directly above one of the detectors in a network of identical detectors will have more power in that detector than any other. If one of the other detectors in the network has more power, then it is less likely that this trigger is a true GW source. Candidate GW signals with the expected ratio of signal power in each detector will have a higher coherent SNR.

This qualitative description of coherence is made rigorous in the following sections up to 4.3.3. In section 4.3.5 we discuss the *null SNR*, another coherent statistic that is useful for identifying background triggers. The rest of this section is about the PyGRB workflow and how to use coherent statistics to confidently detect GWs associated with GRBs.

4.3.1 A Coherent Search for Coalescing Binary Systems

• Show where the A's come from (don't know how to do this yet)

4.3.2 Binary Coalescence Waveform

PyGRB searches for CBC signals with circular orbits and non-spinning components. These waveforms depend on 9 parameters: The two component masses M_1 and M_2 , the sky location (θ, ϕ) , the distance D, the coalescence time t_0 , the orientation ι , the polarisaton angle ψ , and the coalescence phase ϕ_0 . We can reduce to seven parameters if we assume the sky position is known. Of these, the distance, binary inclination, polarisation, and coalescence phase affect the phase and amplitude of the waveform and not the signal morphology. That is, the waveform can be written as

$$h_{+}(t) = \mathcal{A}^{1}h_{0}(t) + \mathcal{A}^{3}h_{\pi/2}(t)$$
(4.1)

$$h_{\times}(t) = \mathcal{A}^2 h_0(t) + \mathcal{A}^4 h_{\pi/2}(t)$$
 (4.2)

where $h_0(t)$ and $h_{\pi/2}(t)$ describe the two phases of the waveform, are usually assumed to be orthogonal, and depend only on the component masses. The amplitudes \mathcal{A}^i are given by

$$\mathcal{A}^{1} = \frac{D_0}{D} \left(\frac{(1 + \cos^2 \iota)}{2} \cos 2\phi_0 \cos 2\psi - \cos \iota \sin 2\phi_0 \sin 2\psi \right) \tag{4.3}$$

$$\mathcal{A}^2 = \frac{D_0}{D} \left(\frac{(1 + \cos^2 \iota)}{2} \cos 2\phi_0 \sin 2\psi + \cos \iota \sin 2\phi_0 \cos 2\psi \right) \tag{4.4}$$

$$\mathcal{A}^3 = -\frac{D_0}{D} \left(\frac{(1+\cos^2 \iota)}{2} \sin 2\phi_0 \cos 2\psi + \cos \iota \cos 2\phi_0 \sin 2\psi \right) \tag{4.5}$$

$$\mathcal{A}^{4} = \frac{D_{0}}{D} \left(-\frac{(1 + \cos^{2} \iota)}{2} \sin 2\phi_{0} \sin 2\psi + \cos \iota \cos 2\phi_{0} \cos 2\psi \right)$$
(4.6)

where D_0 is used to scale the amplitude of the waveforms. For any \mathcal{A}^{μ} we can invert equations 4.3-4.6 to obtain the physical parameters up to a reflection symmetry of the system.

The response of GW detector X to signal $h_{+,\times}$ is given by

$$h^{X}(t) = F_{+}^{X} h_{+}(t^{X}) + F_{\times}^{X} h_{\times}(t^{X})$$
(4.7)

where $F_{+,\times}^X$ is the detector antenna response to the each polarisation and the time of arrival t^X at detector X depends on the sky location of the source and the time of arrival at the fiducial location, e.g. the Earth's center. Combining 4.7 with 4.1 and 4.2 we find

$$h^X(t) = \mathcal{A}^{\mu} h_{\mu}^X(t) \tag{4.8}$$

where the h_{μ}^{X} are given by

$$h_1^X = F_+^X h_0(t^X) (4.9)$$

$$h_2^X = F_{\times}^X h_0(t^X) \tag{4.10}$$

$$h_3^X = F_+^X h_{\pi/2}(t^X) (4.11)$$

$$h_4^X = F_{\times}^X h_{\pi/2}(t^X)$$
 (4.12)

4.3.3 Coherent SNR

The matched-filter between a GW waveform h and the above detector data s is an inner product given by

$$(s|h) = 4\operatorname{Re} \int_0^\infty \frac{\tilde{s}^X(f) \cdot [\tilde{h}^X(f)]^*}{S_h^X(f)}$$
(4.13)

where S_h^X is the noise power spectral density (PSD) for detector X, defined by

$$\langle \tilde{n}^X(f)[\tilde{n}^X(f')]^* \rangle = \delta(f - f')S_h^X(f) . \tag{4.14}$$

Assuming the detector noise is Gaussian, we can work out the likelihood ratio of the signal h being in the data

$$\Lambda(h) = \frac{P(s|h)}{P(s|0)} = \frac{e^{-(s-h|s-h)/2}}{e^{(s|s)}} . \tag{4.15}$$

It is more convenient to work with the log likelihood

$$\log \Lambda = (s|h) - \frac{1}{2}(h|h)$$
 (4.16)

We define the multidetector inner product to be a sum over the d individual detector inner products

$$(\mathbf{a}|\mathbf{b}) = \sum_{X=1}^{d} (a^X|b^X)$$
 (4.17)

The multidetector log-likelihood then becomes

$$\log \Lambda = (\mathbf{s}|\mathbf{h}) - \frac{1}{2}(\mathbf{h}|\mathbf{h}) . \tag{4.18}$$

Using 4.7 we can rewrite the log-likelihood as

$$\ln \Lambda = \left[\mathcal{A}^{\mu}(\mathbf{s}|\mathbf{h}_{\mu}) - \frac{1}{2} \mathcal{A}^{\mu} \mathcal{M}_{\mu\nu} \mathcal{A}^{\nu} \right]$$
 (4.19)

where

$$\mathcal{M}_{\mu\nu} = (\mathbf{h}_{\mu}|\mathbf{h}_{\nu}) . \tag{4.20}$$

The values for \mathcal{A}^{μ} for which the log-likelihood 4.19 is maximal are given by

$$\hat{A}^{\mu} = \mathcal{M}^{\mu\nu}(\mathbf{s}|\mathbf{h}_{\nu}) \tag{4.21}$$

where $\mathcal{M}^{\mu\nu}$ is the inverse of $\mathcal{M}_{\mu\nu}$. We define the *coherent SNR* to be

$$\rho_{\rm coh}^2 = 2 \ln \Lambda |_{\rm max} = (\mathbf{s} | \mathbf{h}_{\mu}) \mathcal{M}^{\mu\nu}(\mathbf{s} | \mathbf{h}_{\nu}) . \tag{4.22}$$

The \mathcal{A}^{μ} terms have been maximised over, leaving only three of our original seven waveform parameters to search over.

4.3.4 Comparison to Coincident Search

In this section we will put the coherent SNR into a form such that it is easy to compare to the coincident SNR. The coincident SNR is defined to be the quadrature sum of the matched filter of the individual detectors

$$\rho_{\text{coinc}}^2 = \sum_{X,Y} \sum_{i=0,\pi/2} \left(s^X \left| \frac{h_i}{\sigma^X} \right) \left[\delta^{XY} \right] \left(s^Y \left| \frac{h_i}{\sigma^Y} \right) \right. \tag{4.23}$$

We can make 4.22 more easily comparable to 4.23 by making \mathcal{M} diagonal. To do this, first note that CBC signals spend a large number of cycles in the sensitive frequency range of the detectors. This causes the 0 and $\frac{\pi}{2}$ phases of the waveform to be approximately orthogonal. The slow frequency evolution means that the two phases have roughly equal amplitude. Hence we find

$$(h_0^X | h_{\pi/2}^X) \approx 0,$$
 (4.24)

$$(h_{\pi/2}^X|h_{\pi/2}^X) \approx (h_0^X|h_0^X) = (\sigma^X)^2. \tag{4.25}$$

Using these, we see that \mathcal{M} simplifies to

$$\mathcal{M} = \begin{bmatrix} A & C & 0 & 0 \\ C & B & 0 & 0 \\ 0 & 0 & A & C \\ 0 & 0 & C & B \end{bmatrix}$$

with

$$A = \sum_{X} (\sigma^{X} F_{+}^{X})^{2} \tag{4.26}$$

$$B = \sum_{X} (\sigma^X F_X^X)^2 \tag{4.27}$$

$$C = \sum_{X} (\sigma^X F_+^X)(\sigma^X F_\times^X) . \tag{4.28}$$

We can simplify further by rotating the detector frame to the *Dominant Polarization Frame*. In this frame, the antenna response functions to the plus and cross polarisation are orthogonal, and hence C=0, making \mathcal{M} diagonal. As we included a polarisation angle between the equatorial and radiation frame (χ) and the radiation and source frame (ψ) , we can rotate our network frame without placing further constraints on the system. In particular, we can rotate through an angle χ^{DP} such that C=0. This gives us the following antenna response functions

$$F_{+}^{\text{DP},X} = F_{+}^{X} \cos 2\chi^{\text{DP}} + F_{\times}^{X} \sin 2\chi^{\text{DP}}$$
 (4.29)

$$F_{\times}^{\text{DP},X} = -F_{+}^{X} \sin 2\chi^{\text{DP}} + F_{\times}^{X} \cos 2\chi^{\text{DP}}$$
 (4.30)

Using these antenna response functions in (4.28) and solving for χ^{DP} , we find

$$\chi^{\rm DP} = \frac{1}{4} \arctan \left(\frac{2 \sum_{X} (\sigma^{X} F_{+}^{X}) (\sigma^{X} F_{\times}^{X})}{\sum_{X} \left[(\sigma^{X} F_{+}^{X})^{2} - (\sigma^{X} F_{\times}^{X})^{2} \right]} \right) . \tag{4.31}$$

This does not uniquely define the dominant polarisation frame, so we also require the network to be more sensitive to the plus polarisation

$$|F_{\perp}^{\text{DP,X}}| \ge |F_{\vee}^{\text{DP,X}}|$$
 (4.32)

In what follows, we assume we are in the dominant polarisation frame and drop the DP superscript. Inverting \mathcal{M} and using (4.22), we see the coherent SNR in the dominant polarisation frame is

$$\rho_{\text{coh}}^{2} = \frac{(\mathbf{s}|\mathbf{F}_{+}\mathbf{h}_{0})^{2} + (\mathbf{s}|\mathbf{F}_{+}\mathbf{h}_{\pi/2})^{2}}{(\mathbf{F}_{+}\mathbf{h}_{0}|\mathbf{F}_{+}\mathbf{h}_{0})^{2}} + \frac{(\mathbf{s}|\mathbf{F}_{\times}\mathbf{h}_{0})^{2} + (\mathbf{s}|\mathbf{F}_{\times}\mathbf{h}_{\pi/2})^{2}}{(\mathbf{F}_{\times}\mathbf{h}_{0}|\mathbf{F}_{\times}\mathbf{h}_{0})^{2}} . \tag{4.33}$$

We can rewrite this as

$$\rho_{\text{coh}}^2 = \sum_{X,Y} \sum_{i=0,\pi/2} \left(s^X \left| \frac{h_i}{\sigma^X} \right) \left[f_+^X f_+^Y + f_\times^X f_\times^Y \right] \left(s^Y \left| \frac{h_i}{\sigma^Y} \right) \right)$$
(4.34)

where we define the orthonormal vectors

$$f_{+,\times}^X = \frac{\sigma^X F_{+,\times}^X}{\sqrt{\sum_Y (\sigma^Y F_{+,\times}^Y)^2}}$$
 (4.35)

We can now compare the coherent SNR as given in (4.34) to the coincident SNR given by (4.23). The coincident SNR is the quadrature sum of all the energy in each detector. In the space spanned by individual detector SNRs, where each trigger is represented by a point in this space, it is the distance from the origin to the trigger. It is the total energy of that trigger.

The coherent SNR is a projection of the total energy of a trigger onto the subspace spanned by f_+ and f_\times . This subspace is the space consistent with a gravitational wave signal from the given sky location and with the PSD of the detectors at the given time. Orthogonal to this space is the space inconsistent with a signal. Noise contributes energy to all components of the coincident SNR, and so projecting it onto the signal space will reduce its amplitude. The energy due to a GW signal lays entirely in the signal plane, so the projection does not change the distance to the point representing the trigger. In the case of a real GW detection, the coherent and coincident SNR are approximately the same as most of the energy of the trigger is due to the GW, though the coherent SNR will be slightly lower as some energy will be due to noise, and so is removed.

Another way to think of the difference between coherent and coincident SNR is in terms of the number of degrees of freedom. The coincident SNR has noise contributions from the phase and amplitude measurements in each detector, resulting in 2N degrees of freedom, where N is the number of detectors. From (4.33) we can see that the coherent SNR has four degrees of freedom: the 0 and $\pi/2$ phases of the plus and cross polarisation amplitudes of the gravitational wave. In the case where we have a non-degenerate¹ two detector network, then both the coincident and coherent SNR have four degrees of freedom. In this case, we can show that the coherent and coincident SNRs are identical by calculating $f_+^X f_+^Y + f_\times^X f_\times^Y = \delta^{XY}$.

4.3.5 Null SNR

Gravitational waves have two polarisations. Therefore we can fully describe the signal with two non-aligned detectors. Adding additional detectors to our network allows us to construct additional data streams that have the GW signal removed. We can easily construct a such a data stream by subtracting the coherent SNR (the energy consistent with a GW signal) from the coincident SNR (the total energy measured by the detectors). Any energy left from this is associated with noise, and we call this the $null\ SNR$

$$\rho_N^2 = \rho_{\text{coinc}}^2 - \rho_{\text{coh}}^2 = \sum_{X,Y} \sum_{i=0,\pi/2} \left(s^X \middle| \frac{h_i}{\sigma^X} \right) [N^{XY}] \left(s^Y \middle| \frac{h_i}{\sigma^Y} \right)$$
(4.36)

where

$$N^{XY} = \delta^{XY} - f_{+}^{X} f_{+}^{Y} - f_{\times}^{X} f_{\times}^{Y} . {(4.37)}$$

¹i.e. sensitive to both plus and cross polarisations.

Noise triggers are incoherent, and so for noise we expect $\rho_{\text{coinc}}^2 \gg \rho_{\text{coh}}^2$. Thus the null SNR is high in this case. For a true GW signal, most of the energy will be coherent, and so we expect $\rho_{\text{coinc}}^2 \simeq \rho_{\text{coh}}^2$. Thus, the null SNR is expected to be close to zero for a true GW signal.

The null SNR is similar to *null streams*, which are used by burst searches (see section 5.2). The null stream in a burst search is made to remove a GW signal of any morphology.

4.3.6 The PyGRB Workflow

We have seen several statistics that can indicate whether a trigger is a GW signal or noise. We apply cuts on these statistics, discarding any triggers that fail the cuts. This is especially important for the coherent search as the coherent statistics are slow to calculate and additional to all the statistics used in the coincident search. To speed up the coherent search, we discard as many triggers as possible at the earliest possible time in the workflow. In this section we will discuss the order of the workflow and the cuts made on each statistic.

The analysis begins with matched filtering the individual detector data. Using the known sky location of the GRB and the fact that gravitational waves travel at the speed of light, we timeshift the data from each detector so that a GW will be coincident in each detector. A list of triggers is formed for each detector by keeping only times when the individual detector SNR is greater than four, the rest of the data is discarded. We then discard any trigger that is not coincident in at least two detectors in the network.

The coincident SNR is then easy to calculate, and any trigger with a coincident SNR below six is discarded. The coherent SNR is then calculated for the remaining triggers. This reduces the SNR of noise, but not signals, and so we apply the same threshold of six to the coherent SNR.

Once we have the coincident and coherent SNR, the null SNR is easy to calculate. The null SNR of a trigger increases with the energy of the trigger for both noise and GW signals, so a very loud GW signal can still have a high null SNR. For this reason, we increase the null SNR threshold with the coherent SNR for loud triggers but keep the null SNR threshold is fixed for quiet triggers, as can be seen in fig.4.1. To be precise, we keep triggers meeting either of the following criteria:

$$\rho_N \le 5.25 \text{ and } \rho_{\text{coh}} \le 20$$
(4.38)

or

$$\rho_N \le \frac{\rho_{\text{coh}}}{5} + 5.25 \text{ and } \rho_{\text{coh}} > 20 .$$
(4.39)

The χ^2 tests (described in 4.2.2) are computationally expensive to calculate, so we calculate these only on those triggers that survive all the other tests. These tests

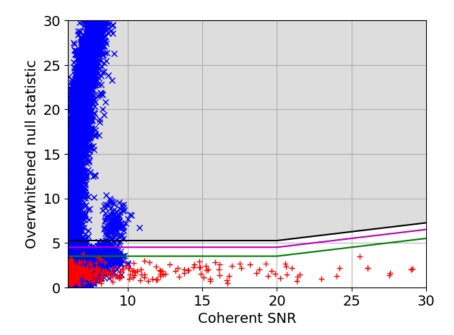


Figure 4.1: **Null Stat Cut.** Here we plot the coherent SNR against the null SNR. The blue crosses are background triggers. The red pluses are signal injections. The black line is the veto line, with all triggers in the shaded region above the line being discarded. The green line indicates the expected SNR for optimally oriented injections. The magenta and cyan lines show 1 and 2 sigma errors on the green line.

are the same as used in the coincident search. In particular, they are not coherent but applied to the individual detector triggers.

- chi2 tests Is this final? No coherent chi2 tests? Do we need to explain why we are no longer using coherent chi2 tests?
- Go through plot caption with Steve

4.3.7 Reweighted SNR

After all the cuts have been made, the remaining triggers are then reweighted twice. First they are reweighted according to their χ^2 values, according to (4.40), then they are reweighted again according to their null SNR, according to (4.41).

Note that this reweighting only down-weights triggers with a high χ^2 or a high null SNR. The reweighting should not significantly alter the network SNR of a GW signal, but will reduce the SNR of noise. The reweighted SNR is detection statistic used by PyGRB.

4.3.8 Searching for Face on Signals

The opening angle of the jet from a short GRB is expected to be $< 30^{\circ}$, emitted perpendicular to the orbital plane. In a similar way to how the coherent SNR accounts for sky position and PSD of each detector to better reject noise, we can build a new detection statistic that takes into account that we expect a true GW signal to have an orbital inclination of ι 0 or ι π with respect to the observer.

We can see from equations (4.3) to (4.6) that the GW amplitude terms depend linearly on $\cos \iota$ and $(1 + \cos^2 \iota)/2$. For $0^{\circ} \le \iota < 30^{\circ}$, these are equal to within 1%. Thus, for ι 0, we can use the approximation $\cos \iota \approx (1 + \cos^2 \iota)/2$. For ι π , the two terms are approximately equal in magnitude, but with opposite signs.

By defining

$$\tilde{D} = \frac{D}{\cos \iota} \text{ and } \chi_{l,r} = \phi_0 \pm \psi$$
 (4.42)

we can rewrite the GW amplitudes (4.3) to (4.6) for $\iota \approx 0$ as

$$\mathcal{A}^1 \approx \mathcal{A}^4 \approx -\frac{D_0}{\tilde{D}}\cos 2\chi_l \equiv \mathcal{B}_1 \tag{4.43}$$

$$\mathcal{A}^2 \approx -\mathcal{A}^3 \approx \frac{D_0}{\tilde{D}} \sin 2\chi_l \equiv \mathcal{B}_2 \ .$$
 (4.44)

We see that now there are only two amplitude factors, \mathcal{B}_1 and \mathcal{B}_2 . Therefore, the GW is circularly polarised. Similar results can be derived for the $\iota = \pi$ case.

Substituting these into (4.19) and working in the dominant polarisation frame², we find

$$\ln \Lambda = \mathcal{B}_1(\mathbf{s}|\mathbf{F}_+\mathbf{h}_0 + \mathbf{F}_\times\mathbf{h}_{\pi/2}) + \mathcal{B}_2(\mathbf{s}|\mathbf{F}_\times\mathbf{h}_0 - \mathbf{F}_+\mathbf{h}_{\pi/2})$$
(4.45)

$$-\frac{1}{2}[\mathcal{B}_1^2 + \mathcal{B}_2^2][(\mathbf{F}_+\mathbf{h}_0|\mathbf{F}_+\mathbf{h}_0) + (\mathbf{F}_\times\mathbf{h}_0|\mathbf{F}_\times\mathbf{h}_0)]. \tag{4.46}$$

Maximising the log likelihood with respect to \mathcal{B}_1 and \mathcal{B}_2 , we find

$$\rho_{\text{coh}}^2 = 2 \ln \lambda_{\text{max}} = \frac{\alpha^2 + \beta^2}{(\mathbf{F}_+ \mathbf{h}_0 | \mathbf{F}_+ \mathbf{h}_0) + (\mathbf{F}_\times \mathbf{h}_0 | \mathbf{F}_\times \mathbf{h}_0)}$$
(4.47)

where

$$\alpha = (\mathbf{s}|\mathbf{F}_{+}\mathbf{h}_{0} + \mathbf{F}_{\times}\mathbf{h}_{\pi/2}) \tag{4.48}$$

$$\beta = (\mathbf{s}|\mathbf{F}_{\times}\mathbf{h}_0 - \mathbf{F}_{+}\mathbf{h}_{\pi/2}) \ . \tag{4.49}$$

In the ι 180° case, the SNR takes the same form, but with

$$\alpha = (\mathbf{s}|\mathbf{F}_{+}\mathbf{h}_{0} - \mathbf{F}_{\times}\mathbf{h}_{\pi/2}) \tag{4.50}$$

$$\beta = (\mathbf{s}|\mathbf{F}_{\times}\mathbf{h}_0 + \mathbf{F}_{+}\mathbf{h}_{\pi/2}) \ . \tag{4.51}$$

²And recalling that $(h_{\pi/2}^X|h_{\pi/2}^X) \approx (h_0^X|h_0^X)$ for CBC signals.

We now have a detection statistic with only two degrees of freedom, as opposed to the four for the coherent SNR and 2N for the coincident SNR in the case of an N detector network. This gives us an additional null stream, and allows us to have a null stream even when analysing data from just two detectors. The circularly polarised coherent SNR allows for a 3% increase in range for a given FAP when compared to the generic coherent SNR. This corresponds to a 10% increase in the rate of GW signals detected.

4.3.9 Searching over a Sky Patch

The sky position of GRBs cannot be determined exactly. The uncertainty in sky position depends on which satellite detected the GRB. The BAT instrument on Swift (see section 3.2) can localise to within 1-4 arcminutes [citation in Andrew's paper]. This is a small enough uncertainty that the GW search can use a single sky point [cite Andrew's paper?]. Other GRB detectors cannot provide as precise localisation. The GBM on the Fermi satellite often provides a location with a roughly circular uncertainty with a radius of several degrees. The IPN uses triangluation to determine sky position, so depending on the number of satellites in the IPN that make the detection, sky localisation and vary from under a square degree to thousands of square degrees.

It is important to have approximately the correct sky position for a coherent search. If the sky position is incorrect, then the detector antenna patterns will be incorrect for the GRB, hurting the search sensitivity. More crucially, with the wrong sky position we will calculate the wrong time delay between detectors in the network and make it impossible to correctly find coincident triggers between detectors.

To search over a sky patch we fill the patch with concentric circles of points separated by δa , such that each ring has $2\pi n/\delta a$ points with n=0 being the central point of the patch. These concentric circles extend to cover 3σ confidence region. The value of δa depends on the depends on the timing uncertainty we are willing to accept. In practice, we use a timing uncertainty of $\delta t=0.5$ ms, which will limit the lost SNR to <5% [cite andrew's paper]. Searching over the sky grid is done after the computationally dominant step of matched filtering, so that searching over the sky grid does not significantly hurt analysis speed [Need some analyses to back this up].

It is interesting to note that in a two detector network, there is a ring of sky points that give the same time delay between the detectors. In this case, only the antenna response of the detectors change with sky position, but the antenna response functions drop out of the coherent SNR in the two detector case³. This would allow for us to analyse a greatly reduced set of sky points. Unfortunately, when we limit

³This is because in the two detector case, both the coherent and coincident SNR have four degrees of freedom. Thus, any observed amplitude and phase is consistent with an astrophysical signal.

our search to looking for circularly polarised GWs, as discussed in section 4.3.8, then the antenna patterns again affect the detection statistic, and we must search over the sky grid.

• Include image of sky grid

4.3.10 Event Significance

Knowing the reweighted SNR of a trigger does not help us know if that trigger is a GW signal or not unless we know the false alarm probability (FAP) of that trigger. The FAP depends on many factors. For example, poor data quality at the time being analysed can lead to more high SNR triggers due to noise. Or the detectors may not be sensitive to the sky position of the GRB, which makes it less likely that a high SNR trigger is a GW signal. In this section we outline the process used to calculate the significance of an event.

As detector noise is not stationary, we calculate the FAP by using the data near the time of the GRB. A GW, if it is there, is assumed to be in the window starting five seconds before the GRB time and ending one second after the GRB time. This window would allow the detection of GWs from most GW models associated with short GRBs. We call this 6s window the *on-source* window. The loudest event, with the loudest template, in the on-source is taken as our trigger.

To evaluate the p-value of this trigger, we analyse approximately⁴ one hour of data around the on-source, called the off-source data. This data is assumed to not contain a GW signal. This data is divided up into as many 6s segments as possible, so that if we have $T_{\rm off}$ seconds of data then we have $N = T_{\rm off}/T_{\rm on}$ segments, where $T_{\rm on}$ is the length of the on-source. These segments are called the off-trials. Using the same criteria as the for the on-source, we find the loudest trigger in each off-trial. As each off-trial is assumed to only contain noise, the FAP of the on-source trigger is the fraction of off-trials that have a louder trigger than the on-source, i.e. the probability that noise would produce a trigger with a reweighted SNR louder than the on-source.

This process alone is only capable of achieving a FAP of 10^{-3} [Cite Andrew's paper] as there is not enough off-trials to claim a lower FAP. This is low enough to reject a signal hypothesis, but not low enough to claim a detection, which requires a FAP of less than 10^{-5} [Cite Andrew again]. One way to get around this would be to use a longer off-source, but as detector noise is not stationary, it is possible that this extra off-source data would have different properties to the time of the GRB. It would also significantly increase the computational time required to complete the analysis.

⁴The amount of data used before and after the on-source varies depending on the data available. For example, sometimes a GRB will happen less than an hour before (after) an interferometer loses (acquires) lock.

To find a lower FAP, the off-source data from each detector is time-shifted relative to the other detectors in the network and the data is then reanalysed. The timeshifts are longer than the light travel time between the detectors in the network and also longer than the duration of a typical glitch (less than one second), so that triggers in the time shifted data are unlikely to appear coherently.

There are two types of time shifting. The first is call short slides and the other long slides. The difference between them stems from the fact that PyGRB analyses data in segments of 128s [Is this still true after the rewrite]. Short slides refers to when only the data within a segment is time shifted. To be precise, in a HLV analysis, the Hanford data is not shifted, the Livingston data is shifted in increments of 6s, and the Virgo data is shifted in increments of 12 seconds. Short slides do not require the data to be matched filtered again, as the time shift happens after the filtering step, so only the network statistics must be recalculated. This means short slides are computationally cheap and can allow for a FAP as low as 10^{-4} . Long slides refers to when the segments themselves are shifted relative to one another. This requires the matched filtering to be redone, as the matched filter time series is not saved when moving to the next segment for analysis. This makes long slides much more computationally expensive than short slides. However, short slides can be done on top of the long slides, so that just 10 long slides are needed to show that a trigger has a FAP $< 10^{-5}$.

It should be noted that our treatment of time slides assumes that each combination of time shifted data is statistically independent of the others. This is not true, as each combination of time shifted data is still the same data being analysed.

- check segment length
- Are we still doing time slides the same way?

4.3.11 Calculating Search Sensitivity

It is useful to estimate the sensitivity of our search to GWs around the time of an analysed GRB. This allows us to know the range to which we are sensitive to GWs in the direction of the GRB, and so rule out the possibility that the GRB was within that distance and producing GWs.

To do this, we inject simulated signals into the off-source data and see if PyGRB can find them. The injected signals are CBC waveforms drawn from three populations: BNS mergers, NS-BH systems with spins aligned with the angular momentum, and NS-BH systems with spins generically aligned. The injections are drawn from an astrophysically motivated distribution of distances, component masses and spins, and binary inclination. The NS mass distribution is a normal distribution with mean $1.4 \rm M_{\odot}$ and standard deviation of $0.2 \rm M_{\odot}$ [refs in O2 GRB paper] restricted to the $1 \rm M_{\odot} - 3 \rm M_{\odot}$ range [refs in O2 GRB paper]. The NS spins magnitude is

restricted to be less than the fastest observed pulsar spin, i.e. ≤ 0.4 [refs in O2 GRB paper]. The BH masses are drawn from a normal distribution of mean $10 \mathrm{M}_{\odot}$ with standard deviation of $6 \mathrm{M}_{\odot}$, restricted to the range $3 \mathrm{M}_{\odot} - 15 \mathrm{M}_{\odot}$. The BH spin magnitudes are ≤ 0.98 , motivated by X-ray observations [refs in O2 GRB paper]. The injections are limited to have binary inclinations or $0^{\circ} - 30^{\circ}$ or $150^{\circ} - 180^{\circ}$, as the opening angle for a CBC powered GRB is expected to be less than 30° .

We quantify the sensitivity of the search using the 50% and 90% exclusion distance. This is the distance at which 50% and 90% of the injected signals are recovered with a greater reweighted SNR than the most significant trigger in the on-source.

4.4 O2 PyGRB Search

- GRB sample
 - Fermi, swift, IPN, others?
 - Need enough science quality data for off source
 - Can analyse single detector data
- Template bank used
- No new detections (just BNS) state BNS p-value
- median exclusion distances

4.5 Improvements to PyGRB

- Made faster matched filter engine and post processing
- Re-optimised the cuts and reweighted snr
- Ran on GRB with bigger sky grid than was previously possible or ran 170817 to be 5 sigma

4.6 Future Plans

- More detectors strengthens the case for coherent search
- Hierarchical search, using coincident search to find location
- Include astrophysical priors in the detection statistic

Chapter 5

Machine Learning for GW Astronomy

5.1 Introduction

The Laser Interferometer Gravitational Wave Observatory (LIGO) and Virgo collaborations operate ground-based Gravitational Wave (GW) detectors. They have detected signals of astrophysical origin, including the merger of a Binary Neutron Star (BNS) system and multiple Binary Black Hole (BBH) merger signals. As these sources have a known signal morphology, they can be found using a highly sensitive matched-filter search, as discussed in chapter 4. But some of the most interesting possible sources, such as core collapse supernova, do not have a known waveform morphology. For this reason, it is important to develop unmodelled searches as well.

In unmodelled searches we look for coherence between the data streams of multiple interferometers. We will consider the case of a Burst search where the sky position of the candidate source is known. This allows us to calculate the relative time of arrival in each detector and the relative signal power in each detector as well.

In this chapter, we will first discuss how an existing targeted Burst search, called *X-pipeline*, searches for Gravitational Wave Bursts (GWBs). We will then see how we can improve this pipeline by using *Multivariate Analysis* (MVA) to apply cuts and rank triggers.

5.2 Xpipeline

Xpipeline is a data analysis pipeline for the Burst search. It begins by using the known sky location of the GRB to time-shift the data from each detector so that the GW arrives simultaneously in each data stream. Xpipeline then makes coherent data-streams from the individual detector data. There are two types of coherent data-streams: signal streams, which increase the power of a GW signals, and null

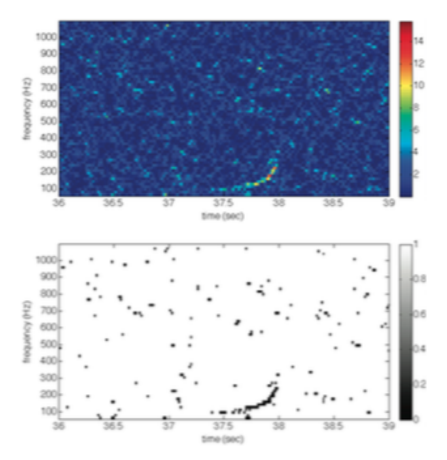


Figure 5.1: **Xpipeline Time-Frequency Map** This figure shows a time-frequency map from Xpipeline for a $1.4 - 10M_{\odot}$ NSBH merger. The top figure shows the E_{+} energy and the bottom figure shows the top 1% of pixels.

streams, which reduce the power of true GW signals but not noise. Triggers are groups of neighbouring pixels that are above threshold in a time-frequency map of a signal stream (see fig.5.1). Xpipeline then removes triggers based on cuts of the various network data streams (see fig.5.2). The position of these cuts is set to give the best performance on a subset of the triggers that are used for tuning. For the MVA pipeline, the position of these cuts is instead set by a machine learning algorithm (as we will see in 5.3).

In this section, we will formulate two of the coherent data-streams generated by Xpipeline, as illustrative examples. The first is a signal-stream, called the *standard likelihood*. We will then use the standard likelihood to create a null-stream.

5.2.1 Burst Search Background

Suppose we have a network of D detectors. A gravitational wave, described by $h_+(t)$ and $h_\times(t)$, passes through the Earth from direction $\hat{\Omega}$. We describe the sensitivity of detector $\alpha \in \{1, ..., D\}$ to the plus and cross polarizations using the *Antenna*

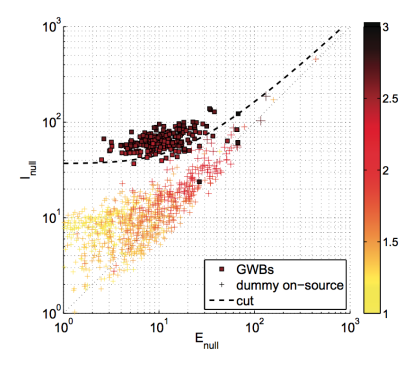


Figure 5.2: **Xpipeline Cut** This figure shows Xpipeline making a cut to eliminate many spurious signals.

Response Functions, denoted $F_{\alpha}^{+}(\hat{\Omega})$ and $F_{\alpha}^{\times}(\hat{\Omega})$. The position of detector α is denoted by \mathbf{r}_{α} and the noise in this detector is given by $n_{\alpha}(t)$. The detector output $d_{\alpha}(t)$ is then given by

$$d_{\alpha}(t + \Delta t_{\alpha}(\hat{\Omega})) = F_{\alpha}^{+}(\hat{\Omega})h_{+}(t) + F_{\alpha}^{\times}(\hat{\Omega})h_{\times}(t) + n_{\alpha}(t + \Delta t_{\alpha}(\hat{\Omega})) . \tag{5.1}$$

Here Δt_{α} is the time taken for the GW to reach the detector from some arbitrary reference point¹ \mathbf{r}_0

$$\Delta t_{\alpha}(\hat{\Omega}) = \frac{1}{c} (\mathbf{r}_0 - \mathbf{r}_{\alpha}) \cdot \hat{\Omega} . \qquad (5.2)$$

From now on we will suppress explicit mention of the reference point \mathbf{r}_0 or the time delay Δt_{α} on the understanding that detector outputs need to be time-shifted by an appropriate amount.

In reality, detector outputs are not continuous but sampled discretely. The discrete Fourier transform $\tilde{x}[k]$ of the time series x[j], and its inverse, are given by

$$\tilde{x}[k] = \sum_{j=0}^{N-1} x[j]e^{-i2\pi jk/N}, \qquad x[i] = \frac{1}{N} \sum_{j=0}^{N-1} \tilde{x}[k]e^{i2\pi jk/N} . \tag{5.3}$$

For sampling rate is f_s and N data points in the time domain, we convert continuous

¹The center of the Earth is a fairly intuitive choice for a worldwide detector network.

to discrete notation by using

$$x(t) \to x[j] \tag{5.4}$$

$$\tilde{x}(f) \to f^{-1}\tilde{x}[k]$$
 (5.5)

$$\int dt \to f_s^{-1} \sum_j \tag{5.6}$$

$$\int df \to f_s N^{-1} \sum_k \tag{5.7}$$

$$\delta(t - t') \to f_s \delta_{jj'} \tag{5.8}$$

$$\delta(f - f') \to N f_s^{-1} \delta_{kk'} . \tag{5.9}$$

For example, the one-sided noise spectral density for a detector with noise n(t) can be written in continuous form as

$$\langle \tilde{n}^*(f)\tilde{n}(f')\rangle = \delta(f - f')\frac{1}{2}S_n(f)$$
(5.10)

where the angle brackets indicate an average over the noise. In the discrete notation listed above, this becomes

$$\langle \tilde{n}_{\alpha}^{*}[k]\tilde{n}_{\beta}[k']\rangle = \frac{N}{2}\delta_{\alpha\beta}\delta_{kk'}S_{\alpha}[k] . \qquad (5.11)$$

We will be working with the noise-spectrum-weighted quantities, defined by

$$\tilde{d}_{w\alpha}[k] = \frac{\tilde{d}_{\alpha}[k]}{\sqrt{\frac{N}{2}S_{\alpha}[k]}}$$
(5.12)

$$\tilde{n}_{w\alpha}[k] = \frac{\tilde{n}_{\alpha}[k]}{\sqrt{\frac{N}{2}S_{\alpha}[k]}}$$
(5.13)

$$F_{w\alpha}^{+,\times}(\hat{\Omega},k) = \frac{F_{\alpha}^{+,\times}(\hat{\Omega})}{\sqrt{\frac{N}{2}S_{\alpha}[k]}}$$
(5.14)

The normalisation of the whitened data is

$$\langle \tilde{n}_{\alpha}^*[k]\tilde{n}_{\beta}[k']\rangle = \delta_{\alpha\beta}\delta_{kk'} . \tag{5.15}$$

In vector notation, we can write 5.1 as

$$\tilde{\mathbf{d}} = \mathbf{F}\tilde{\mathbf{h}} + \tilde{\mathbf{n}} \tag{5.16}$$

where $\mathbf{F} = [\mathbf{F}^+ \ \mathbf{F}^{\times}]$ and $\tilde{\mathbf{h}} = [\tilde{h}_+ \ \tilde{h}_{\times}]^T$.

5.2.2 Standard Likelihood

Assume that the noise in our detectors is Gaussian. As a gravitational wave $\tilde{\mathbf{h}}$ passes through the detector from a known direction, the probability of attaining whitened detector output $\tilde{\mathbf{d}}$ in one time-frequency pixel is given by

$$P(\tilde{\mathbf{d}}|\tilde{\mathbf{h}}) = \frac{1}{(2\pi)^{D/2}} \exp\left[-\frac{1}{2} \left|\tilde{\mathbf{d}} - \mathbf{F}\tilde{\mathbf{h}}\right|^2\right] . \tag{5.17}$$

For a set $\{\tilde{\mathbf{d}}\}$ of N_p time-frequency pixels, we have

$$P(\lbrace \tilde{\mathbf{d}} \rbrace | \lbrace \tilde{\mathbf{h}} \rbrace) = \frac{1}{(2\pi)^{N_p D/2}} \exp \left[-\frac{1}{2} \sum_{k} \left| \tilde{\mathbf{d}}[k] - \mathbf{F}[k] \tilde{\mathbf{h}}[k] \right|^2 \right] . \tag{5.18}$$

By comparing this value to the probability that the detector produces this output in the absence of any GW signal, we can calculate a likelihood of the signal being a real GW signal.

The Likelihood Ratio L is the log of the probability that the detector network will have output $\tilde{\mathbf{d}}$ in the presence of GW $\tilde{\mathbf{h}}$ divided by the probability of obtaining the same output in the absence of a gravitational wave ($\tilde{\mathbf{h}} = 0$)

$$L = \ln \frac{P(\{\tilde{\mathbf{d}}\}|\{\tilde{\mathbf{h}}\})}{P(\{\tilde{\mathbf{d}}\}|\{0\})} = \frac{1}{2} \sum_{k} \left[\left| \tilde{\mathbf{d}} \right|^{2} - \left| \tilde{\mathbf{d}} - \mathbf{F}\tilde{\mathbf{h}} \right|^{2} \right] . \tag{5.19}$$

For the above analysis to be applied, we would need to know the waveform $\tilde{\mathbf{h}}$ in advance. For GRBs and other unmodelled searches, this is not possible. One way to handle this problem is to fit the waveform in each time-frequency pixel to the data in such a way as to maximise the likelihood ratio. Hence we have

$$0 = \frac{\partial L}{\partial \tilde{\mathbf{h}}} \bigg|_{\tilde{\mathbf{h}} = \tilde{\mathbf{h}}_{\text{max}}} \tag{5.20}$$

Solving this, we find

$$\tilde{\mathbf{h}}_{\mathbf{max}} = (\mathbf{F}^{\dagger} \mathbf{F})^{-1} \mathbf{F}^{\dagger} \tilde{\mathbf{d}}$$
 (5.21)

where the superscript dagger † denotes the conjugate transpose.

Calculating the likelihood ratio for $\tilde{\mathbf{h}}_{\mathbf{max}}$ gives us the Standard Likelihood E_{SL}

$$E_{\rm SL} = 2L(\tilde{\mathbf{h}}_{\rm max}) = \sum_{k} \tilde{\mathbf{d}}^{\dagger} \mathbf{P}^{\rm GW} \tilde{\mathbf{d}}$$
 (5.22)

where

$$\mathbf{P}^{\mathrm{GW}} \equiv \mathbf{F} (\mathbf{F}^{\dagger} \mathbf{F})^{-1} \mathbf{F}^{\dagger} . \tag{5.23}$$

We can see from equation 5.1 that the contribution made to the data output by a passing GW from fixed sky location is restricted to the subspace spanned by \mathbf{F}_{+}

and \mathbf{F}_{\times} . Therefore the energy in this subspace is the energy that is consistent with a GW from a given sky location. We can show that \mathbf{P}^{GW} is a projection operator, projecting the data into this same subspace. The standard likelihood maximises the energy in this subspace, and so is the maximum energy contained in the whitened data that is consistent with a GW from the given sky location.² This is an example of the coherent signal-streams that Xpipeline uses.

5.2.3 Null Energy

We can use the standard likelihood to find a null-stream. First consider the *total* energy, given by

$$E_{\text{tot}} = \sum_{k} |\tilde{\mathbf{d}}|^2 \ . \tag{5.24}$$

This is an incoherent statistic as it contains only auto-correlation terms, and no cross-correlation terms. If we subtract the standard likelihood from the total energy, we obtain the *null energy*

$$E_{\text{null}} \equiv E_{\text{tot}} - E_{\text{SL}} = \sum_{k} \tilde{\mathbf{d}}^{\dagger} \mathbf{P}^{\text{null}} \tilde{\mathbf{d}} .$$
 (5.25)

This is the energy that is inconsistent with a GW from given sky location, and must therefore be associated with noise. This is the minimum amount of energy in our whitened data that is inconsistent with our GW. It is an example null-stream used by Xpipeline.

This shows one of the key advantages of coherent analysis. If we analysed our data incoherently, we would be working with just the total energy. But using coherent methods, we can project the whitened data into the subspace spanned by \mathbf{F}^+ and \mathbf{F}^\times , thus removing some fraction of the noise without removing any of the signal. The drawback is that if the sky position is not known in advance, then the analysis needs to be repeated for a set of directions that span the entire sky ($\gtrsim 10^3$ directions), each with different antenna response functions \mathbf{F}^+ and \mathbf{F}^\times . This will slow down analysis and increase the false alarm probability.

5.2.4 Incoherent Energy and Background Rejection

The diagonal elements of 5.25 are auto-correlation terms, and the other elements are cross-correlation terms. The auto-correlation part of the null energy are called the *incoherent energy*, and denoted by

$$I_{\text{null}} = \sum_{k} \sum_{\alpha} P_{\alpha\alpha}^{\text{null}} |\tilde{d}_{\alpha}|^2 . \qquad (5.26)$$

²In practice, some of this energy will be due to noise. We say that the energy due to noise is *inconsistent* with a GW signal. The rest of the energy must be due to the GW, so we say it is *consistent* with the GW.

Background triggers are typically not correlated between the different detectors of the network, so the cross-correlation terms are small relative to the auto-correlation terms. This means that for glitches, we have

$$E_{\text{null}} \approx I_{\text{null}}$$
 (5.27)

Compare this to the case of a GW signal. This will be correlated between the detectors. By construction, the energy of the correlated energy does not appear in the null-stream. Therefore, in the presence of a strong GW, the incoherent energy is much larger than the null energy

$$E_{\text{null}} \ll I_{\text{null}}$$
 (5.28)

Using 5.27 and 5.28, we see that the ratio of E_{null} and I_{null} is very different in the case of a glitch as opposed to a GW signal. We can use this to make the following cut to remove noise triggers from our sample

$$I_{\text{null}}/E_{\text{null}} > C$$
 (5.29)

for some constant C > 1. This test does not work as well for small amplitude glitches, where the statistical fluctuations can lead to E_{null} being smaller than I_{null} . For this reason, Xpipeline varies C with the size of the trigger. The precise position of the cut is set to maximise performance on a set of injections.

5.3 Machine Learning

We have seen how Xpipeline makes cuts on coherent statistics to distinguish between noise and GW signals. You can see from 5.29 that the cut only uses two statistics, the null energy and the incoherent energy. Xpipeline generates many coherent statistics that could used in conjunction to make a more powerful cut. In this section we discuss how to use Machine Learning to make achieve this. The software we use is the Toolkit for Multivariate Analysis package in the ROOT data analysis framework.

5.3.1 Supervised Machine Learning

The type of Machine learning we use is called *supervised machine learning*. Supervised machine learning algorithms are trained on data which has already been classified. They can then be shown a new, not-classified data point and determine the appropriate classification. Supervised machine learning requires data to be in a particular format (see table 5.1). It is a list of *events*, each with a *label* and corresponding *attributes*. The machine learning algorithm builds a *classifier* that can determine the label of an event when given the event's attributes. In the case of our GW search, the events are the triggers returned by Xpipeline (see section 5.2). The

Label	loghbayesiancirc	standard	circenergy	circinc
Background	12.3128	58.3523	44.7196	24.9015
Background	12.0349	67.5344	41.7045	22.4848
Signal	18.2145	59.8136	53.3320	22.0601
Signal	43.7113	123.9194	118.9774	43.9234
Signal	6422.1467	14426.9124	14167.2933	4991.7876

Table 5.1: An example of MVA training data. Each event has a label and several attributes.

labels are *signal* or *background*, and the attributes are the values of the signal and null data streams for those triggers as well as some statistics describing the trigger, e.g. peak frequency, trigger duration. A full list of the attributes used, together with a short description of the attribute can be seen in table 5.2. The signal triggers are generated by injecting signal waveforms into the data. Background triggers are triggers that do not coincide with an injected signal.

The signal and background trigger sets are each divided into two subsets: The training set and the testing set. The training set is used to build the classifier, while the testing set is used to measure the accuracy of the trained classifier. If the classifier performs much better on the training set than on the testing set, then the classifier is overtrained. This means that the classifier has learned exactly the properties of signals and noise in the training set, rather than learning the general properties of signals and noise. When an overtrained classifier is used on a different data set with different noise properties (such as the testing set), it will perform poorly.

Coherent Energy	Description	
loghbayesiancirc	A likelihood ratio based on Bayesian methods, for the hypothesis of a circularly polarised GW vs Gaussian noise	
$E_{ m max}$	The maximum energy in the whitened data that is consistent with a GW from a given sky location.	
$E_{ m null}$	The minimum amount of energy in the whitened data that is inconsistent with a GW from a given sky location. Given by $E_{\rm tot}-E_{\rm max}$.	
$I_{ m null}$	The sum of the autocorrelation terms of $E_{ m null}$.	
$E_{ m circ}$	The maximum energy in the whitened data consistent with a circularly polarised GW from a given sky location.	
$I_{ m circ}$	The sum of the autocorrelation terms of the $E_{\rm circ}$.	
$E_{ m circnull}$	The energy in the whitened data that is inconsistent with a circularly polarised GW from a given sky location. Given by $E_{\rm tot}-E_{\rm circ}$.	
$I_{ m circnull}$	The sum of the autocorrelation terms of E_{circnull} .	
$E_{ m H1}$	The energy in the H1 interferometer.	
$E_{ m L1}$	The energy in the L1 interferometer.	
$E_{ m V1}$	The energy in the V1 interferometer.	
number of pixels	The number of pixels in the cluster.	
Duration	Time duration of the trigger.	
Bandwidth	The frequency range spanned by the trigger.	
Power law		

Table 5.2: The attributes used by the machine learning classifier.

5.3.2 Boosted Decision Trees

A Decision Tree is a simple type of classifier. It is a flowchart of true/false statements about a trigger's attributes to determine the correct label. For example, consider the decision tree shown in fig.5.3. Here the attributes are labeled as the components of a vector x. We start at the top node and work downwards. If the node is true then we follow the branch to the left and if not then we follow the branch to the right. We then consider the statement at the end of whichever branch we follow. We continue this process until we reach a leaf node, which has no branches and contains a final classification for the trigger.

We can improve the performance of the classifier by using an *ensemble* (or *forest*) of trees. This means training multiple distinct trees, and each tree independently classifies the trigger. The final classification of each trigger is a normalised sum of the outputs of each tree, with +1 corresponding to signal, and -1 corresponding to background. This leads to different regions of the parameter space having different MVA scores, as can be seen in fig.5.4. The higher the score, the more likely an event is to be a signal.

To train a decision tree, we need methods to find the variable to cut on and the position of the cut for each decision node, and the label in each leaf node that best discerns between signal and background. Each of these values is set by brute force: Trying each possible combination of cuts and labels to get the best performance on the training events. Ensemble methods work best when each classifier in the ensemble is independent of the others, so training every tree on the same events is not going to give optimal results. For this reason each tree is trained on some subset of the training set. We could pick events at random to form these subsets, but a more powerful method is to use *Adaptive Boosting*.

When using adaptive boosting, each event in the training set is given a probability that it will be selected to train the next tree. Initially each event in the training set has the same probability of being selected. Then after each tree is trained the probability of each event being selected to train the next tree is updated such that misclassified events are more likely to be included in the training set for the next tree. The misclassified events have their probability of being selected for training the next tree updated by the boost weight, given by

$$\alpha = \frac{1 - \text{err}}{\text{err}} \tag{5.30}$$

where err is the misclassification rate. The weights are then renormalised.

Note that the boost weight is greater than one for³ err< 1/2, and that as the error gets smaller, the boost weight increases. The effect of this is that each new tree is more likely to be trained on the events that are most difficult to classify.

³The misclassification rate is always less than half when the labels only take two possible values as the algorithms we use are always better than chance alone.

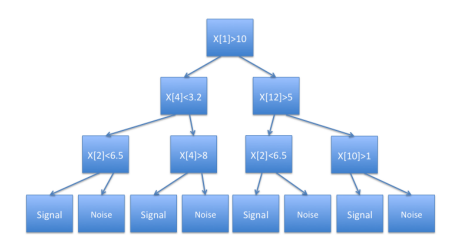


Figure 5.3: A decision tree example To determine if a trigger is a signal or noise event the tree makes a series of cuts on the attributes x[N]. If the inequality in a node is true, then the next node is the branch to the left. Otherwise the next node is the one to the right.

The ensemble output is also changed, so that it is weighted rather than being a simple sum. If the output of the *i*th tree is given by $h_i(\mathbf{x})$, with \mathbf{x} being the event attributes, then the ensemble output is given by

$$H(\mathbf{x}) = \frac{1}{N} \sum_{i=0}^{N} \ln(\alpha_i) h_i(\mathbf{x})$$
 (5.31)

where N is the number of tree in the ensemble. Small values indicate background events, while large values indicate a signal.

5.3.3 Data Preprocessing

Extra data preprocessing is required for the training set. Suppose a small amplitude signal injection is added close to a large amplitude glitch in the data. This trigger will be labeled as a signal, due to the injection, but the properties of the trigger will resemble a glitch, as the glitch has a much larger amplitude than the signal injection. It is essentially a background trigger labeled as a signal. This has two affects: It reduces our ability to detect real signals by making the properties of signals harder for the algorithm to learn, and it can lead to background triggers being misclassified as a Gravitational Wave signal.

For these reasons, it is important to make sure that signal injections do not overlap in time or frequency with background triggers. To prevent this from happening

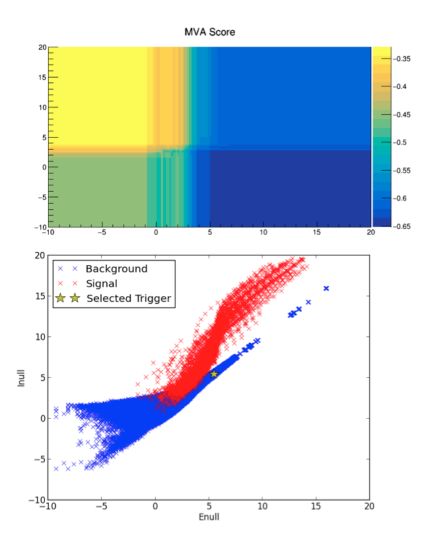


Figure 5.4: Visualising the Classifier In the bottom plot you can see the value for log(Enull) and log(Inull) for all the signal and background training data used to build the classifier. We chose one of these events at random (indicated by the star) and varied Enull and Inull to see how it changed the MVA score, indicated by the colour in the top plot. As you can see, increasing Inull and decreasing Enull leads to the event being more likely to be classed as a signal. This is akin to the xpipeline cut shown in fig.5.2.

we clean the data. This means finding all of the triggers in the data for the smallest signal injection scale. The injected waveforms are too small to be detected so all of these triggers must be background. We then increase the injection amplitude and look for triggers again. Any triggers that overlap in time or frequency with the noise triggers are then identified and removed from the from the signal set. This description of cleaning may need to change if we change the cleaning codes.

We must also not include injections in our signal set that are too small to be detected. If we do then we would again be including triggers labeled as signal that have the properties of a noise trigger, even if there is no glitch present. For this reason we apply a threshold on the amplitude of the signal set, so any injection below that amplitude is removed. The level of the threshold is set experimentally; If it is too low then we will increase the chance of a false positive and hurt our sensitivity to real signals, but if it is too high then we will limit the classifiers ability to detect low amplitude signals.

We also use Generalised Clustering. This is a change in the way Xpipeline defines a trigger. Without using generalised clustering, triggers are groups of neighbouring pixels in the time-frequency maps such as in fig.5.1. Generalised clustering allows these pixels to be separated by a user-specified number of pixels. This can boost the signal power of long triggers, as long triggers tend to contain breaks in their time-frequency maps that causes Xpipeline to list a single long trigger as multiple short triggers. The downside to using generalised clustering is that it can cause noise triggers to be grouped together as well, boosting the power of noise. This then causes our sensitivity to short triggers to be reduced slightly. Experimenting with generalised clustering in both the standard Xpipeline analysis and the MVA analysis suggests that the benefits outweigh the costs, with an improvement of between 7% and 50% for long inspiral and adi waveforms at the cost of a decrease of 3% to 9% for short sine-gaussian waveforms.

As XTMVA is to be used for the unmodelled search, it is also important to ensure that the search can find waveforms that are not in the training set. There are several tools that we use to achieve this. The first is to limit the amount of information the classifier is given about the waveform morphology. The classifier cannot know the precise morphology of any waveform because the only attributes that the classifier trains on is the time, peak-frequency, and various measurements of the coherent and null energy between the detectors in the network. This forces the classifier to use the coherence of the triggers to make a classification, rather than the waveform morphology.

There is still a possibility that the classifier will become too specialised to the waveforms in the training set, as the classifier is given peak frequency data and certain waveform morphologies may have particular characteristics than become apparent from the coherent and null streams. For this reason we also trained the

classifier on a variety of different waveforms. The training set included long and short waveforms, and a variety of different morphologies. Some of the signals are astrophysically motivated, such as compact binary coalescence signals, while some are artificial, such as the white noise burst.

The final tool we use to ensure the classifier is sensitive to generic waveforms is to test the classifier on waveforms that are not included in the training set. If the classifier can find waveforms not in the training set, then we can be reasonably confident that it is sensitive to generic waveforms. We also try removing certain waveforms from the training set to test the robustness of the classifier. This will lead to a drop in sensitivity for that waveform, but if the drop in sensitivity is small, then we can be confident that the classifier is robust.

5.3.4 Optimisation and Validation

With any machine learning algorithm, there are hyperparameters that must be tuned. The number of trees in the ensemble, the maximum depth of the trees, and many more hyperparameters must be chosen. We optimise the hyperparameters by repeatedly running an example analysis with different hyperparameters, trying to increase the number of injected signals we could recover from the testing set.

Setting our hyperparameters using the testing set can cause data leakage. Data leakage is when data from outside the training set is used to build a classifier. As we tune our classifier on the testing set, it is possible that we will implicitly tune our classifier to only work well on this training and testing set. This is very similar to overtraining discussed in section 5.3. To avoid this, once we have tuned the hyperparameters on a single GRB, we test the classifier on several other GRBs. If the performance drops significantly on these other GRB analyses, then we have had data leakage and we need to retune our classifier. This process of testing on previously unused data is called validation. If there is evidence of data leakage, then we must retune our classifier and validate it again, this time using a different (previously unused) GRB so as to avoid data leakage from the validation GRB.

Optimisation is a somewhat cyclical process, as once we have changed one hyperparameter, we must then go back and test that other parameters do not now need changing. But optimisation is not pure guess work, and in the rest of this section we will see the strategy we used to optimise our classifier. We first discuss optimising the training data to use. As mentioned in section 5.3.3, there are several choices to make regarding what data is used for training, such as setting the amplitude threshold. We then discuss optimising the BDT classifier itself.

Training Set

Many of the hyperparameters mentioned in section 5.3.3 had to be optimised. Consider the amplitude threshold applied to triggers before they are included in the

training set. If this is too low then noise triggers contaminate the signal training set, but if it is too high then we limit the sensitivity of our classifier to only higher amplitude waveforms.

We also optimised the waveforms that are included in the training set, whether to use cleaning or not, and whether to use generalised clustering. Write more on this once we've done it.

- Had to optimise the threshold as well
- Training waveforms
- cleaning or not
- Gen clustering or not

BDT Parameters

There are many hyperparameters that need to be set for a BDT analysis. In this section we discuss some of these hyperameters and the method we used to optimise them for our analysis.

We began by setting values for *NTrees*, the number of trees in the ensemble, and the learning rate, discussed in section 5.3.2. These two parameters are set first to ensure the machine learning algorithm will converge in a reasonable amount of time. Setting the learning rate too low causes the classifier to take longer to converge. Setting the learning rate too high can cause the classifier to never converge. Similarly, using too many trees takes too long for the classifier to finish training, but too few and the training will terminate before the algorithm has converged. Setting these first ensures that we have a classifier that gives sensible results in a reasonable amount of time. While optimising, we set these values slightly high, to ensure that our classifier converges and does so quickly while we optimise our other parameters. Once the other parameters are set, we again optimise NTrees and the learning rate, increasing the number of trees and decreasing the learning rate to ensure the algorithm reaches its optimum performance, even if it increases the time taken for training.

We now tune the tree-specific parameters. Unlike the number of trees and the learning rate which are primarily tuned to ensure the classifier will converge in a reasonable time, these parameters are set to ensure that the classifier is accurate but does not overtrain. Overtraining can happen when the trees are allowed to make cuts that are too fine, carving out regions of parameter-space around anomalous events in the training set rather than finding cuts that generalise beyond the training data. The way to prevent this is to limit how fine the cuts made by the decision trees are allowed to be, while allowing cuts that are fine enough to pick out the general features of signal and background events in our data. There are several hyperparameters we can set to do this, which must all be tuned.

The first of these is the maximum depth of the trees, which is the maximum number of cuts a tree can make before it reaches a leaf node. Each cut divides the parameter-space into ever smaller regions which it labels as background or signal. Setting the maximum number of cuts too low will therefore cause the classifier to be too coarse in dividing up the parameter-space, resulting in poor accuracy. Increasing the maximum depth allows the classifier to pick out smaller features in the parameter-space. If the maximum depth is too high then the classifier will overtrain; dividing the parameter-space into precisely the regions that work for the training set and losing generality. As we are using adaptive boosting, it is recommended cite TMVA userguide to use trees with fewer cuts. For this reason we tried values from 2-16, finding that for our problem a maximum depth of 8 was optimal.

A related hyperparameter is the minimum number of events that we allow in a leaf node. If we allow the training algorithm to have any number of events in a leaf node, then it will occasionally find cuts that results in a small number of events in one or more of the leaf nodes. The again allows for the carving out of anomalous events in the training set rather than finding cuts that can generalise beyond the training set. Conversely, setting the minimum number of events allowed in the leaf nodes to be too high does not allow the classifier to pick out the key features in the data. We used a grid search over the values 100-1600 for the minimum number of events per leaf node and found the optimal value to be 400.

The final hyperparameter we set is the number of cuts that the training algorithm scans over to find the best cut. When the classifier is training it searches for the best way to cut the parameter-space into a signal subspace and a background subspace. To do this we can try every possible cut on every parameter. This can be very slow and lead to overtraining. To speed up our analysis and reduce the chance of overtraining, we can choose the number of cuts to try on each parameter. For example, we may decide to use 10 cuts for each parameter. In this case the algorithm will tune the cut on a parameter which ranges from 0-100 by trying cuts at 0,10,20,... and selecting the best of these cuts. To tune the number of cuts we wanted to use, we tried values from 10-160 as well as allowing the classifier to try every possible cut. We found that 80 cuts was optimal for our purposes.

• This can definitely be phrased better

Result of Optimisation

- Show nice plots of improved sensitivity due to optimisation
- Show improvement when compared to Xpipeline