Nonparametric Transient Classification using Adaptive Wavelets

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ABSTRACT
Classifying transients based on the multi band light curves is a challenging but crucial problem in the era of GAIA and LSST since the sheer volume of transients will make spectroscopic classification unfeasible. Here we present a nonparametric classifier that uses the transient’s light curve measurements to predict its class given training data. It implements two novel components: the first is the use of the BAGIDIS wavelet methodology – a method of characterizing functional data using hierarchical wavelet coefficients. The second novelty is the introduction of a ranked probability classifier on the wavelet coefficients which handles both the variable size of measurement errors (heteroscedasticity) of the data in addition to the potential non-representativity of the training set. The ranked classifier is simple and quick to implement while a major advantage of the BAGIDIS wavelets is that they are translation invariant, hence they do not need the light curves to be aligned to extract features. Further, the BAGIDIS methodology is nonparametric so it can be used for blind searches for new objects. We demonstrate the effectiveness of our ranked wavelet classifier against the well-tested Supernova Photometric Classification Challenge (SNPCC) dataset in which the challenge is to correctly classify light curves as belonging to Type Ia or non-Ia supernovae. We train our ranked probability classifier on the spectroscopically-confirmed subsample (which is not representative) and show that it gives good results for all supernova with observed light curve timespans greater than 100 days (roughly 55% of the dataset). For such data, we obtain a Ia efficiency (recall) of 80.5% and a purity (precision) of 82.4% yielding a highly competitive score of 0.49 averaged over all redshifts whilst implementing a truly “model-blind” approach to supernova classification. The classifier compares favourably to standard algorithms such as k-Nearest Neighbours (kNN) and Support Vector Machines (SVM), which obtain scores of 0.45 and 0.41 respectively. Consequently this approach may be particularly suitable for the classification of astronomical transients in the era of large synoptic sky surveys.

Key words: supernova, photometric classification, wavelets, splines, BAGIDIS, ranked probability classifier.

1 INTRODUCTION
A major shift occurring in astronomy is the dawn of wide-field synoptic surveys such as GAIA and LSST that will sweep up in their nets vast numbers of objects which then require automated sifting and classification; a trend that is pushing astronomy rapidly into the arena of machine learning.

The lure of machine learning lies in its potential to exploit large quantities of data to provide significant insights despite limited prior understanding by training on known data. This, of course, can be problematic if the training data is not representative, i.e. is systematically different in some way from the main test data that needs to be classified. In astronomical applications the training data is typically created by taking spectra to classify the class of a transient.
Unfortunately spectra are expensive to take, especially at high redshift, meaning that training data are unlikely to be representative but rather biased towards brighter objects and lower redshifts and hence not representative.

A good example of the impact of this lack of representativeness is provided by the 2010 photometric supernova classification challenge (Kessler et al. 2010) in which many groups competed using a number of different methods for separating the light curves of Type Ia supernovae (SNIa) from those of types Ib/c and II (labelled collectively as “non-Ia” or “nIa”). The training data was designed to mimic current spectroscopic followup strategies which yielded a highly biased and non-representative dataset that had a much higher Ia/nIa ratio than in the test dataset. This lack of representativity wrecked havoc with many of the classification algorithms used for the challenge. In Newling et al. (2011) it was found that a training set of 50 representative supernovae gave the same performance as a training set with approximately 1000 unrepresentative supernovae.

While it is possible in principle to produce a representative training data, it is difficult to do so. At the faintest end even if one schedules spectroscopic followup the resulting spectrum may be too noisy to yield a clear identification. Even more tricky, one needs the training dataset to be representative for all classes of objects to be classified, which for general wide-field synoptic surveys such as GAIA, SKYMAPPER and LSST will include AGN, variables stars as well as the various sub-classes of supernovae. It is likely that a truly representative training sample for such surveys will be difficult to assemble. Therefore, it is important to develop features and classification algorithms that are relatively robust to the expected types of non-representativity in future astronomical surveys.

In this paper we search for methods that (1) are not significantly affected by the non-representativity of the training set, (2) that do not require the light curves to be aligned, (3) do not require any redshift estimate for the object and (4) make minimal assumptions about the form of the light curves ("non-parametric"). Photometric supernovae classification has traditionally been performed by comparing the supernova’s light curve measurements to the light curve templates for the various supernova classes. The supernova class whose template most closely matched the candidate supernova’s measurements would be the predicted class. However such methods are very sensitive to the error in the templates with small deviations from the true templates causing severe drops in classification performance. Recently, cosmologically-blind approaches have been proposed. These typically fit parametric models to the light curves and use the resulting parameter estimates as features that one may use to build a classifier. These approaches have been shown to work well provided the parametric form of the flux model is consistent with the flux behaviour of the transient objects being studied. As such a parametric approach – such as that followed in Newling et al. (2011) – is not suitable when dealing with astronomical objects whose transient behaviour is poorly understood or when the transient behaviour varies drastically between the classes of the transient object.

Future photometric surveys will not only be detecting supernovae of course, but will also capture other transient astronomical objects – some of whom may represent new classes altogether and which may encode much of the value of the new survey. In such cases, parametric curves for the new classes or sub-classes do not exist and so a general nonparametric method of capturing the transient behaviour is advisable. We advocate using the coefficients of a wavelet projection as a general means of characterizing the behaviour of transient astronomical objects. See Fuga (2004) for a thorough introduction to wavelets. Wavelets are a popular method of representing curves which simultaneously show localised sharp features (peaks, troughs, ...) and smooth parts. They combine the property of any non-parametric curve estimation method to take advantage of neighbouring points being more likely to have values close to each other, with the wavelet’s built-in localisation: in fact the curve representation is done by adding a series of localised basis functions (e.g. of compact support). Having this property, wavelets tend to outperform principal components analysis as means of dimension reduction when applied to functional data with localised sharp transients. We thus apply a wavelet analysis to characterize the light curves, subsequently fitting various classifiers to the resulting dataset. We also advocate the use of a ranked probability classifier as it is relatively robust to nonrepresentative training sets and can deal with heteroskedastic data. With heteroskedastic data, the astronomical measurements errors do not all share the same variance. Both these issues are relatively common in astronomical datasets.

Previous work on classifying astronomical transients from their light-curves includes the extensive work on the 300 million light curves from the Catalina Real-Time Transient Survey Philip et al. 2012; Donalek et al. 2013; Graham et al. 2013; Djorgovski et al. 2014; Faraway et al. 2014 and the GAIA detector Long et al. 2012; Blagorodnova et al. 2014). Wavelets have been used in regression with astronomical spectra in Machado et al. (2013) to estimate redshifts. The focus of the Machado et al. (2013) paper was to minimize catastrophic failures in the redshift estimation process.

We use wavelets to characterize functional astronomical data within the Supernova Photometric Classification Challenge (SNPCC) dataset. The SNPCC dataset simulates flux measurements under realistic observing conditions. Consequently, the time of first detection varies greatly with some supernovae only detected well after peak-brightness. This source of heterogeneity is a problem for traditional functional data characterization as most characterization methods would consider curves that have the same shape but which are shifted with respect to each other to be very different. Indeed, dealing with shifted curves is a problem with both template fitting methods as well as the cosmology-blind parametric methods. The standard solution to the varying locations is to align the flux measurements for the various supernovae so that time zero corresponds to the estimated time at which the various supernovae reach peak brightness. This can be computationally intensive to implement and may not be appropriate for other forms of astronomical transient objects. We thus propose using the BAGIDIS methodology to characterize the curves. BAGIDIS is a hierarchical wavelet procedure that is robust to shifts in the positions of curves and which can also be readily applied to non-dyadic datasets – that is datasets where the curves are of arbitrary length (Timmermans et al. 2013; Timmermans & von Sachs 2014).

In section 2 we describe a method for extracting features
from functional data. The method can be readily extended to deal with heteroskedastic data as well as functional data that is not equidistant. In section 3, we propose a classifier that is relatively robust to a non-representative training set. This is followed in section 4 by a description of the SNPPC dataset. The methods in sections 2 and 3 are applied to the SNPPC dataset in section 5 which describes the implementation of the feature extraction and classification procedures while in section 6 we compare and discuss the performances of the various implementations. Finally, we make some concluding remarks in section 7.

2 FEATURE EXTRACTION FOR FUNCTIONAL ASTRONOMICAL DATA

Most astronomical datasets contain examples of what in the statistics literature is termed “functional data” where each observation consists of a set of samples of one (or perhaps several) function(s) with a single argument. For the time being, we shall assume for each of $N$ astronomical objects that there is a single corresponding function $f_i$ that we measure. For the $i$-th astronomical object we obtain $n_i$ measurements of the corresponding function $f_i$ at the argument values $x_{i,1}, x_{i,2}, \ldots, x_{i,n_i}$ which are denoted $y_{i,1}, y_{i,2}, \ldots, y_{i,n_i}$. $y_{i,j} = f_i(x_{i,j}) + \epsilon_{i,j}$ where $i = 1, \ldots, N, j = 1, \ldots, n_i$ (1) with $\epsilon_{i,j}$ denoting the measurement errors, which are supposed to be normally distributed $\epsilon_{i,j} \sim \text{Normal}(0, \sigma_{i,j}^2)$. Examples of functional data within astronomy include astronomical spectra (here $y_{i,j}$ is the measured light intensity of the $i$-th astronomical object at the wavelength $x_{i,j}$) as well as the light curves of variable objects such as supernovae (here $y_{i,j}$ is the measured flux of the $i$-th supernova at time $x_{i,j}$ in a particular colour band).

One application of functional data is in the field of classification. Suppose we have $N$ astronomical objects that are divided into $M$ classes (where $M \ll N$) and that for each object we wish to sample a function $f$ as an aid to classification. For many contexts, it is reasonable to assume that the behaviour of the function $f$ is determined by the astronomical object’s class membership. Let $\tau_i$ represent the class of the $i$-th astronomical object (that is $\tau_i \in \{1, 2, \ldots, M\}$ for $i = 1, \ldots, N$). If class membership uniquely determines the functional form then eqn (1) can be rewritten as:

$y_{i,j} = f_{\tau_i}(x_{i,j}) + \epsilon_{i,j}$ where $i = 1, \ldots, N, j = 1, \ldots, n_i$. (2)

That is we have $M$ unique but unknown functions \(f_{1}, \ldots, f_{M}\) with the total number of independent samplings over these $M$ functions being $N$. Assume for each astronomical object that we have an equal number of measurements of their functions $\{f_{\tau_i} : i = 1, \ldots, N\}$. That is $n_i = n$ for $i = 1, \ldots, N$. The more the functions $\{f_{\tau_i} : i = 1, \ldots, N\}$ are sampled (that is, the larger $n$ becomes), the more information there is of the behaviour of these functions for the various classes. However, if these samples are used directly as input for a classification procedure, increased sampling of the function may paradoxically decrease classification accuracy since the extra information gained from the larger $n$ may not compensate for the increased dimensionality of the input space. Increased dimensionality makes inference difficult for a number of reasons: firstly as the dimensionality of an input space increases, the expected distance between two randomly chosen sampled functions, $(y_{1,1}, y_{1,2}, \ldots, y_{1,n})$ and $(y_{j,1}, y_{j,2}, \ldots, y_{j,n})$ increases. That is, data becomes more sparse as the dimensionality of the input space increases. Furthermore, the complexity of the classification rule often increases with the dimensionality of the input space thus dramatically increasing the variance of the estimated classification boundary.

Consequently, it is worthwhile to seek to reduce the dimensionality of the input space whilst preserving as much information within the dataset as possible. The traditional and most popular method for dimension reduction is principal components analysis (PCA). Here, the $N$ observations of the $n$ correlated samples are transformed into a set of (at most) $n$ linearly uncorrelated principal components. Principal components analysis (PCA) has been used to successfully compress stellar spectra by a factor of over 30 (Bailer-Jones et al. 1998). The reduced dataset was then fed into a neural network for classification. Instead of using PCA for dimension reduction, we instead propose using wavelets for performing dimension reduction. Wavelets are uniquely tailored to represent functional data whose underlying signal consists of a time-varying mixture of frequency components. Such functional datasets are prevalent in the astronomical literature and hence wavelets are a natural tool for analysis of many astronomical datasets. In the next subsection we propose a method of wavelet analysis that can be readily applied to a wide range of functional astronomical data.

2.1 The BAGIDIS Methodology

In our initial discussion of the BAGIDIS methodology we will make two main points that will subsequently be relaxed in forthcoming sections. Firstly, we assume for every astronomical object that there is a corresponding function $f_i$ that we wish to estimate and that all $N$ of these functions are sampled on the same equally-spaced grid i.e. $n_i = n = n_j$ with $(x_{i,1}, x_{i,2}, \ldots, x_{i,n}) = (x_{j,1}, x_{j,2}, \ldots, x_{j,n})$ for $i, j \in \{1, \ldots, N\}$ and also that $x_{i,(k+1)} - x_{i,k} = \Delta x_{i,(l+1)} - x_{j,l}$ where $i, j \in \{1, \ldots, N\}$ and $k, l \in \{1, \ldots, n-1\}$. Secondly, we assume that the measurements error variances are the same i.e. $\sigma_{i,j}^2 = \sigma^2 = \sigma_{j,j}^2$ for $i, i' \in \{1, \ldots, N\}$ and $j, j' \in \{1, \ldots, n\}$. As pointed out in the introduction to this section, we would like to benefit from the ability of wavelet methods to compress the significant information of curves into a small number of relevant features (Antoniadis et al. 2001), used for subsequent (functional) classification.

The higher the compression (i.e. the reduction of the dimensionality) – without losing significant information – the better for the subsequent classifier. However, in this context and in fact more generally, a recurring problem in functional data analysis is to estimate the similarity of various curves. Traditional methods of estimating curve similarity tend to be very sensitive to shifts in the observed curves. As an example, if two curves have exactly the same shape, but one of the curves is shifted by an amount $H$ from the other, then many measures of curve similarity will consider the two aforementioned curves to be very dissimilar. This problem is particularly relevant to astronomical data sets, as e.g. astronomical spectra as well as the SNPPC dataset, to be analysed subsequently here: the onset of a peak in the spectrum or in a light curve can be slightly shifted from one
data set to another without making the two curves necessarily belong to different classes.

Previous approaches have circumvented the aforementioned problem by first aligning the sampled curves. In the case of supernova classification, this alignment is often achieved by first attempting to find the time at which each light curve achieves peak brightness and subsequently shifting the time axis so that time zero corresponds to the time of peak brightness (Newling et al. 2011; Ishida & de Souza 2013). Though such an alignment procedure works well in practice, there are a number of disadvantages to such an approach. Firstly, implementing such a procedure is considerably more difficult if the function has several peaks. This is not the case for the various supernova classes, however one can expect future photometric surveys to find transient objects that have multiple peaks and so aligning the curves may be difficult. Secondly, the alignment procedure could be computationally intensive. Thirdly, many alignment procedures make implicit assumptions about the form of the functional data. Such assumptions may not necessarily be valid for all classes of astronomical objects. We thus choose to implement a comparatively new wavelet procedure – called BAGIDIS (Timmermans et al. 2011) – that explicitly accounts for possible shifts in the curves. Furthermore, unlike traditional wavelet analysis, BAGIDIS does not require the dataset to be dyadic. This is important as the sequence of measurements for the curve will often not be dyadic. Finally, as a key to success, BAGIDIS uses a hierarchical ordering of its wavelet coefficients in order to encode “optimally” the significant features of various, potentially misaligned curves with the purpose of discriminating them by a small number of coefficients. We describe the BAGIDIS methodology in more detail now.

The acronym BAGIDIS stands for BAses GIving DIStances, as basis expansion is at the core of the methodology, together with the introduction of a new distance measure in the coefficient domain of this expansion. The BAGIDIS methodology has been introduced in Timmermans & von Sachs (2010). Timmermans & von Sachs (2014), where significant gains in performing dimension reduction over other methods, such as PCA, have been demonstrated (e.g. in a blood-serum analysis of spectra). Further investigation of its use in nonparametric functional prediction and classification (Ferraty & Vieu 2006) is provided in Timmermans et al. (2011). The following description of the key ideas is general enough for application of the method to any set of curves with one or more localised features (spikes, peaks, troughs, discontinuities,...) such as light curves and spectra, not just those resembling the supernova light curve dataset that we shall subsequently use to demonstrate the methodology which are of relatively short length and which show at most two localised humps.

As a first step of the procedure, if necessary by the length of the data, each curve is decomposed in a set of short series using a sliding window, so that each windowed series should not contain two significant patterns of the same amplitude (in our specific application this was not necessary). Subsequently the (windowed) series is expanded in a particular wavelet basis, with wavelet coefficients called $d_k$, which are referred to as the “Unbalanced Haar Wavelet Basis” (UH-basis$^3$). It is obtained using the “Unbalanced Haar Wavelet Transform” (UH-WT$^4$) proposed by Fryzlewicz (2007). This general version of BAGIDIS – where each curve is decomposed into a set of short series – offers the possibility to localise the analysis along shorter windows. This decomposition prevents otherwise similar features that are not just slightly shifted but are far apart from each other from being treated as the same feature (in case this is undesirable from the nature of the problem at hand).

Haar wavelets are made of level-change indicator functions on a hierarchy of dyadic subintervals of the unit (observation) domain, they are piecewise constant to the left and the right of the level-change point and normalised such that they constitute an $L_2$– orthonormal basis (akin to the Fourier basis but now being localised on short intervals and hence comprising much better compression properties). Unbalanced Haar wavelets overcome the dyadic restriction: the level-change (or breakpoint) $b_k$ can be at any of the observation points of the underlying signal. Still they constitute an orthonormal basis when implemented via the UH-WT: in fact, the set $\{(b_k, d_k)\}_k$ determines the shape of the projected signal uniquely, and induces an interesting property of hierarchy in the resulting UH-basis expansion. The most important level change of the signal, i.e. its location and its amplitude, is encoded in $(b_1, d_1)$, whilst secondary local features are to be found in higher coefficients. An illustration of this along a simple noiseless example made of 9 data points is shown in Fig. 1. As hoped for and observed by looking at the location of the discontinuity points $b$ between positive and negative parts of the wavelets, the first non-constant vectors support the largest discontinuities of the series and encode therefore the highest peak of the series. Subsequent vectors point to smaller level changes while the last few vectors correspond to zones where there is no level change – as indicated by the associated zero coefficient.

In a noisy set-up, the UH-WT coefficients with “ranks” (i.e. indices) $k$ higher than some integer $D$ do not carry any more significant information (but just noise contribution) and should be suppressed from the basis expansion. (The critical rank $D$ is usually small but needs to be user-chosen – ideally in a data-driven way – as depending on the problem at hand.) In such a way, the UH-WT allows for an automatic and unique hierarchical description of each of the curves into a curve-adapted orthonormal basis. The hierarchy makes the resulting bases (UH-basis) comparable to each other, although different, and allows subsequent classification based on a distance of similarity of two measured curves $y^{(1)}$ and $y^{(2)}$ – each of which have been sampled $n$ times – via similarity of their $\{(b_k, d_k)\}_{k=1}^D$ coefficients (where $D \leq n$):

$$d(y^{(1)}, y^{(2)}) = \sum_{k=1}^{D} w_k \left( \lambda |b_k^{(1)} - b_k^{(2)}|^2 + (1 - \lambda)|d_k^{(1)} - d_k^{(2)}|^2 \right)^{1/2}$$

(3)
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Figure 1. Left panel: Illustration of a UH-WT expansion. The left panel consists of three parts denoted I., II., and III. respectively. In the upper part of the left panel (part I.) we plot the series. The corresponding abscissa axis at the very bottom is common for that graph and for the graph of basis vectors (part II.). The main part of the figure (part II.) shows the basis vectors of the Unbalanced Haar wavelet basis that is best suited to the series (UH-WT basis). These vectors are represented rank by rank, as a function of an index along the abscissa axis and are denoted $\psi_0$, ..., $\psi_8$ where the subscript denotes the rank. Dotted horizontal lines indicate the level zero for each rank. Vertically, from top to bottom on the right hand side (part III.), we find the detail coefficients associated with the wavelet expansion. Each coefficient is located next to the corresponding basis vector. Right panel: Representation of the series in the breakpoint-wavelet coefficient (b-d) plane. The same series is plotted in the plane that is defined by the values of its breakpoints and its detail coefficients. Points are numbered according to their rank in the hierarchy.

Here the tuning parameter $\lambda \in [0, 1]$ can be chosen according to the application at hand (or even by supervised learning). In cases where the astronomical curves are shifted with respect to each other yet where we choose to avoid any aligning of the curves, the BAGIDIS procedure is still able to recognize that the curves are the same since the hierarchical ordering of the ranks is performed in a data-driven manner. Hence, by putting all the weight in eqn [8] on the $d_k$’s, two curves that are shifted with respect to each other will be considered identical. With the weight function $w_k$ one has the flexibility to give more weight on more important features (which are encoded in the first coefficients). Again this parameter can be chosen in a supervised way, e.g. by cross-validation.

Selecting the optimum number $D$ of BAGIDIS coefficients for classification involves a tradeoff: the more coefficients we use to represent the dataset, the smaller the information loss. However, more coefficients will cause the classifier to have higher variance. The optimum number of BAGIDIS coefficients to train a classifier can be selected from the data, by e.g. a cross-validation procedure, by choosing the number of BAGIDIS coefficients that minimize the misclassification error rate for the validation sets.

2.2 Estimating the Variances of the BAGIDIS Coefficients

In the previous section a distance measure (given by eqn [8]) was proposed. Note that since BAGIDIS imposes a hierarchical ordering upon its wavelet coefficients, we’d typically expect the weights $w_k$’s to be a monotonically decreasing function of $k$. Timmermans & von Sachs (2010) proposed that the weights be given by:

$$w_k = \frac{\log(n + 1 - k)}{\log(n + 1)}$$

where $n$ is the number of measurements of the function $f$. It was found that this functional form worked well for many datasets. However whatever the functional form chosen, the weights (as shown in eqn [8]) do not depend on the pair of observed curves $(y^{(1)}, y^{(2)})$ for which we wish to calculate a distance measure. This simplified structure is natural if the input is homoskedastic. Indeed, homoskedasticity is a standard assumption for most traditional classifiers. However for astronomical data, there will often be a large level of heterogeneity in the variances of the wavelet coefficients. This heterogeneity stems from the underlying heterogeneity of the measurement process: both the preciseness and the number of measurements collected for the pool of astronomical objects within a database can commonly vary by at least an order of magnitude. This heterogeneity must be accounted for within the distance measure.

Suppose that the variances of the $k$-th BAGIDIS co-
coefficients for the observations $y^{(1)}$ and $y^{(2)}$ are denoted by $\sigma_k^{(1)^2}$ and $\sigma_k^{(2)^2}$ respectively. If these variances are known, then a more appropriate distance measure would be:

$$d(y^{(1)}, y^{(2)}) = \sum_{k=1}^{D} \left[ \frac{w_k}{\sigma_k^{(1)^2} + \sigma_k^{(2)^2}} \left( \lambda |b_k^{(1)} - b_k^{(2)}|^2 \right) + (1 - \lambda)|d_k^{(1)} - d_k^{(2)}|^2 \right]^{1/2}$$

(4)

We thus turn our attention to the estimation of the variances $\sigma_k^{(1)^2}$ and $\sigma_k^{(2)^2}$ of the BAGDIS coefficients.

For astronomical datasets, each measurement on an object often includes a corresponding estimate of the variance of its measurement error. In principal it could be possible to derive an analytical expression for how the variance of the measurement errors filters through to a set of wavelet coefficients. However, such an approach is not feasible for the BAGDIS coefficients as the underlying wavelet coefficients are reordered in order of importance. This re-ordering makes the derivation of an analytical formula for the variances of the BAGDIS coefficients very difficult.

Instead, we wish to estimate the variances by Monte Carlo simulation. Assume the measurement errors for the functions $\{f_i : i = 1, ..., N\}$ are Gaussian with mean zero and known variance. Under this assumption we may use Monte Carlo simulations to estimate the variances of the BAGDIS features. This entails using the distributional information on the measurement errors to resample a new set of equidistant measurements for each function. By subsequently applying the BAGDIS procedure to each simulated set of equidistant measurements, we can investigate how the randomness of the measurement errors feeds through to the BAGDIS coefficients. By simulating several sets of BAGDIS coefficients for each astronomical object, we create a pool of coefficients that we can use to precisely estimate the variances of the BAGDIS coefficients for each astronomical object. This variance estimation procedure will be modified in section 2.3 to allow for non-equidistant measurements. In cases where the resulting simulated BAGDIS coefficients may have outliers, one can use M-estimators instead of the sample variance as an estimator of the true variance. See Chapter 5 of van der Vaart (2000) for more details about the M-estimators.

2.3 Allowing for Non-Equidistant Functional Data

Given a sequence of $n$ equidistant observations, the BAGDIS methodology will return $n$ wavelet coefficients that are ordered according to the importance of the sequence’s level changes. Thus in order to use BAGDIS to compare the similarity between pairs of astronomical objects, we require their corresponding functions to each be sampled on the same equidistant grid. We assumed the existence of such a grid in sections 2.1 and 2.2. However, this assumption is rarely met by astronomical datasets since observing time on instruments are limited and instruments are often subject to adverse observing conditions. Consequently, it is important that we be able to adapt any arbitrary sampling of the function of interest into a format that is usable by the BAGDIS methodology. We propose the utilization of splines to create the required dataset.

Suppose we have $N$ astronomical objects sampled at the points $\{(x_{i,1}, x_{i,2}, ..., x_{i,n_i}) : i = 1, ..., N\}$ and suppose that $x_{i,1} = 0$ and $x_{i,n_i} = T$ for all $i$. That is, for all $N$ astronomical objects we have measurements that span the interval $[0, T]$. Under such conditions, it is possible to fit a spline to each object over the region $[0, T]$. The spline estimates at points along an equidistant grid within $[0, T]$ can subsequently be used as the input for the BAGDIS procedure.

Given a sequence of measurements $\{y_{i,j} : j = 1, 2, ..., n_i\}$ at times $\{x_{i,j} : j = 1, 2, ..., n_i\}$ for the $i$-th object, we wish to find a function $g_i(x)$ that will fit this data well (that is have a small residual sum of squares) whilst also being reasonably smooth. A natural cubic spline – with knots at the points $(x_{i,1}, x_{i,2}, ..., x_{i,n_i})$ – will be the function $g_i(x)$ that minimizes the following objective:

$$\sum_{j=1}^{n_i} (y_{i,j} - g_i(x_{i,j}))^2 + \lambda_i \int_0^T g_i''(x)^2 \, dx.$$  

(5)

The first term is the sum of squared residuals for the cubic spline whilst the second term is a regularization factor that penalizes roughness within $g_i(x)$. $\lambda_i$ represents a tuning parameter for the $i$-th object that controls the trade-off between the fidelity to the data and the roughness of the spline estimate. Since astronomical data is typically heteroskedastic (that is the measurement errors do not all share the same variance), we instead find the spline that minimizes:

$$\sum_{j=1}^{n_i} \frac{1}{\sigma_{i,j}^2} (y_{i,j} - g_i(x_{i,j}))^2 + \lambda_i \int_0^T g_i''(x)^2 \, dx.$$  

(6)

where as before $\sigma_{i,j}^2$ is the variance of the $j$-th measurement error for the $i$-th object. As such, the resulting splines tend to follow the precisely measured fluxes more closely, thus making the method relatively robust to outliers resulting from imprecise measurements. For more details on natural cubic splines, refer to sections 5.2.1 and 5.4 of Hastie et al. (2009).

We choose the value of the tuning parameter $\lambda_i$ for each astronomical object (and hence our cubic spline function) by cross-validation. If we wish each astronomical object to be sampled on an equidistant grid of length $n^*$, we can set $x_{i,1} = 0, x_{i,2} = \Delta, x_{i,3} = 2\Delta, ..., x_{i,n^*} = (n^* - 1)\Delta$ for $i = 1, 2, ..., N$ with the sampled values given by the corresponding spline estimates i.e. $y_{i,1} = g_i(0), y_{i,2} = g_i(\Delta), y_{i,3} = g_i(2\Delta), ..., y_{i,n^*} = g_i((n^* - 1)\Delta)$. This sequence of $n^*$ spline estimates can be taken as input for the BAGDIS procedure. The $'$ on each variable denotes that the variable is part of a data sequence that has been adapted to be equidistant. $\Delta$ is a tuning parameter whose value we choose. Note that if $\Delta$ is too large, the resulting grid will be very coarse and some information on the function of interest will be lost. On the other hand the smaller $\Delta$ is, the more computationally expensive will be the feature extraction. In order to get the variances of the BAGDIS coefficients, we again use Monte Carlo simulation: for each astronomical object, we simulate a set of measurements. For each set of simulated measurements, we can fit a spline using the inverse variance weighting, sample the fits on an equidistant grid and use the resulting sequence as input for the BAGDIS procedure. Fig. 5 shows the resampled splines for the flux measurements of a supernova with the accompanying de-
3 A RANKED PROBABILITY CLASSIFIER

Suppose we have \( N \) astronomical objects whose class \( \tau \) is known (the training set) and an additional \( L \) objects of unknown class (the test set). For astronomical datasets, the training set will often have a different distribution (e.g., lower redshift, lower apparent magnitude) than the test set. This difference in distribution is problematic as many classifiers are sensitive to non-representativity. One approach to making a classifier more robust to non-representativeness is to make pairwise comparisons between the candidate from the test set and each object in the training set. The predicted class for the candidate will then only be a function of the \( K \) closest matches within the training set where \( K \) can be chosen by cross-validation. This subset of the training set are the only objects that influence the predicted class and they will typically be more representative of the test set when the input space is low-dimensional. The most common example is the \( K \) nearest neighbour classifier (kNN). Ishida & de Souza (2013) successfully applied kNN to classify supernova light curves with a highly non-representative set. However, kNN cannot give the class probabilities and does not naturally account for the heteroskedasticity of the dataset. We thus develop a ranked classifier that can address these two shortcomings.

Assume for all \( N + L \) objects that we have obtained a sequence of \( n \) measurements of a function \( f \) (as in eqn 2) on an equidistant grid. Hence the information on the function \( f \) – compressed into a vector \( \mathbf{d} = (d_1, d_2, ..., d_D) \) of BAGIDIS coefficients – can be used to classify the \( L \) test objects of unknown type. The proposed ranked probability classifier works very similarly to kNN. For each candidate object within the test set, we wish to calculate the probability that it’s observed vector \( \mathbf{d_c} \) could have arisen from measurement error if the candidate in reality shared the same underlying function as (say) the \( i \)-th object within the training set. Assuming Gaussianity and independence, the relevant density is given by:

\[
g(\mathbf{d}_c|\mathbf{d}_i) = (2\pi)^{-D/2}|\Sigma_c + \Sigma_i|^{-1/2} \exp\left\{-\frac{1}{2}(\mathbf{d}_c - \mathbf{d}_i)^T(\Sigma_c + \Sigma_i)^{-1}(\mathbf{d}_c - \mathbf{d}_i)\right\}
\]

(7)

where \( \Sigma_c = \text{diag}(\sigma_{1,c}^2, ..., \sigma_{D,c}^2) \) and \( \Sigma_i = \text{diag}(\sigma_{1,i}^2, ..., \sigma_{D,i}^2) \). We wish to calculate this density for all objects in the training set i.e. \( g(\mathbf{d}_c|\mathbf{d}_i) : i = 1, ..., N \). The candidate is then classified as being of the same class as the training set object that yields the highest density value. One can also calculate the approximate probability of the candidate object belonging to a class \( m \) using:

\[
\Pr(\tau_C = m) \approx \frac{g(\mathbf{d}_c|\mathbf{d}_m^*)}{\sum_{i=1}^{M} g(\mathbf{d}_c|\mathbf{d}_i^*)}, \quad m = 1, ..., M
\]

where \( \mathbf{d}_i^* \) is the BAGIDIS vector of the closest matching object within the training set that belongs to class \( i \). Though we are able to calculate the class probabilities, this ability was only possible through assuming the coefficients came from an independent Gaussian distribution. BAGIDIS coefficients often have a Gaussian distribution or can easily be transformed to a Gaussian distribution. Unfortunately the independence assumption is highly unlikely to be met due to the re-ordering of the coefficients. That said, the classifier seems to be relatively robust to the violation of the independent Gaussian assumption.

4 THE SUPERNOVA PHOTOMETRIC CLASSIFICATION CHALLENGE (SNPCC) DATASET

For the purpose of this paper we use the Supernova Photometric Classification Challenge (SNPCC) dataset. The SNPCC dataset consists of simulated light curves for over 18,000 supernovae. Less than 7% of these supernovae are spectroscopically observed. This subset are the supernovae for which the supernova class is known. The challenge was released in January 2010 and consisted of trying to develop a classifier based on the spectroscopic subset (the training set) that would accurately predict the class membership of the rest of the dataset (the test set) using the supernova's light curve data. In particular, the focus is on correctly identifying the type Ia supernova within the test dataset.

For each supernova, we have a series of flux measurements over four frequency bands (G-, R-, I- and Z-bands) together with both the time of the measurement as well as
the corresponding flux measurement errors. Fig. 2 shows the measurements recorded for a randomly selected supernova. These light curve measurements were simulated under a variety of observing conditions. Consequently the datasets are highly variable: the number of measurements taken and the flux measurement error vary greatly from one supernova to the next. Fig. 3 shows the distributions of both the number of flux measurements as well as the observed timespans (elapsed time between the first and the last flux measurement of the supernova) in the G-band for the combined SNPCC dataset. The top panel is clearly bimodal with a high proportion (≈ 40%) of the supernovae having less than 20 flux measurements taken. The number of flux measurements is strongly correlated with the supernova’s time span (ρ ≈ 0.9) with roughly the same proportion of supernovae having an observation time span of less than 100 days. In contrast, for currently running photometric surveys (and indeed future surveys) we expect these proportions to be much smaller. The times at which measurements are taken are irregular and differ between supernovae. These highly variable datasets are not suitable for many traditional methods of analysis that often assume that the data is standardized. It is thus important to find a general method – such as that proposed within section 2 – to extract a standardized set of features for each of the supernova.

With the SNPCC dataset, the training set is also highly unrepresentative of the test set. The non-representativity arises due to the procedure for spectroscopic follow-up. Since spectroscopic followup is expensive, the training set tends to consist of supernova with much higher apparent magnitudes (see Fig. 4). Also since type Ia supernova are generally brighter than non-Ia’s, the proportion of Ia’s within the training set (≈ 70%) is much higher than the proportion of Ia supernovae within the test set (≈ 30%). In addition, the redshifts of the Ia supernova within the test set tends to be higher than that for the training set. In the case of the non-Ia supernovae, the test set will tend to have a pool of supernovae that are either intrinsically dimmer or that are of higher redshift. We thus need to develop a classifier that is not overly sensitive to the non-representativity of the training set. These two aforementioned problems of non-standardized datasets and non-representativity shall be addressed using the methods proposed within sections 2 and 3. We discuss the application of these methods to the SNPCC dataset in the next section.

5 APPLICATION OF THE METHODOLOGY TO THE SNPCC DATASET

5.1 Feature Extraction for the SNPCC dataset

In order to deal with the wide variation in both the number of flux measurements as well as the observed time spans for the supernovae within the SNPCC dataset, we wish to obtain for each supernova a set of flux estimates on an equidistant grid. For the SNPCC analysis, we require flux estimates at the grid values (0 days, 2 days, 4 days, ..., 98 days, 100 days). The more frequently we sample the flux estimates, the less information is lost and the more computationally expensive is the feature extraction procedure. We chose a 2 day sampling frequency as it results in minimal information loss whilst still being computationally feasible. We use cubic splines (section 2.3) to estimate the flux values at the required times, with the weights of each measurement being inversely proportional to its variance (see eqn 6). Fig. 5 demonstrates how the inverse weighting results in more stable spline fits. The regularisation parameter is chosen by cross-validation, with the parameter constrained to lie on the interval 0.45 ≤ λ ≤ 0.9. This restriction assumes that the underlying function that we are sampling is slowly varying (due to the lower bound) and that it is probably nonlinear (due to the upper bound). For the majority of the supernovae, the cross-validated value of λ lies well within the interval.

For each supernova, we have four sequences of 51 measurements. We wish to compress the resulting 204-dimensional space into a smaller subspace so as to reduce
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Figure 4. Density plot of the average measured flux in the R-band for each supernova. The red density plot is for the training set and the blue plot is for the test set. The supernovae within the training set tend to be considerably brighter than the supernovae within the test set.

Figure 5. Observed flux measurements for a randomly chosen Ia supernova over four frequency bands. In all four panels, two cubic splines are fitted to the observations. The dashed lines is a standard cubic spline whilst the solid line represents the cubic spline that results when the flux measurements are weighted inversely by the variance of the measurement error. The latter spline seems to provide the more stable fit.

Figure 6. The top panel shows the R-band measurements for a randomly chosen supernova (black dots with error bars) together with the spline estimates on the equidistant grid (purple triangles). The next five panels show progressively more accurate reconstructions of the original spline estimates using incrementally more BAGIDIS coefficients.

For each sequence, we compress the sequence into $D$ coefficients using the BAGIDIS methodology. The coefficients are hierarchical with the first BAGIDIS coefficient in each sequence storing the sequence mean. Since the training data is brighter than the test data, we ignore the first coefficient (as any decision rule based on the first coefficient is unlikely to generalise well when moving to the test set) and rather compress the original measurements into the 2nd, 3rd, ..., $(D + 1)$-th BAGIDIS coefficients. $D$ is chosen so as to maximise the cross-validated classification accuracy. Fig. 6 shows how the reconstructions of a randomly chosen supernova become more accurate as $D$ increases. Using $D = 5$ coefficients in each of the four colour bands provides over 10-fold compression of the spline estimates whilst also preserving the bulk of the light curve information. Using $t$-distributed stochastic neighbourhood embedding (see van der Maaten & Hinton (2008) for more details), we are able to represent the resulting 20-dimensional BAGIDIS input vector for each supernova within the combined training and test sets using a two-dimensional subspace (see Fig. 7). Reassuringly, we find that the Ia’s and non-Ia’s are very
well separated in this two-dimensional subspace. Hence, the BAGIDIS coefficients are providing excellent discriminatory information for classification.

For each supernova, we thus have four vectors of $D$ BAGIDIS coefficients. We expect the randomness in the measurement process to have fed through to the coefficients and so it is important to also calculate a vector of the standard deviations of the BAGIDIS coefficients as these may be used to aid in classification. These vectors may be calculated using Monte Carlo simulation (see section 2.2). Fig. 5 shows a randomly selected supernova in the R-band together with the 1,000 splines fits to the simulated measurements. The standard deviations for the R-band were used within the classifiers as cross-validation revealed that their use results in the fewest misclassifications. With both the BAGIDIS coefficients and their corresponding standard deviations, we can turn our attention to the classification of each supernova within the test set.

5.2 Application of Selected Classifiers to the Extracted BAGIDIS Coefficients

We apply three classification algorithms to the SNPCC dataset. Two are well-known classification algorithms and the third is the ranked probability classifier proposed in section 3. For each of these classifiers, we need to estimate the classification performance. Kessler et al. [2010] suggested the following score for the SNPCC:

$$S_{\text{FoM-Ia}} = \frac{N_{\text{true-Ia}}}{N_{\text{TOT-Ia}}} \times \frac{N_{\text{true-Ia}}}{N_{\text{false-Ia}}} + \frac{W_{\text{false-Ia}}}{N_{\text{false-Ia}}}$$  \hspace{1cm} (8)

where $N_{\text{true-Ia}}$ is the number of test set supernovae that are correctly classified as Ia's, $N_{\text{false-Ia}}$ is the number of test set supernovae that are incorrectly classified as Ia's, $N_{\text{TOT-Ia}}$ is the total number of Ia's within the test set and $W_{\text{false-Ia}}$ is the false-Ia tag penalty factor. The first fraction represents the Ia efficiency – the proportion of Ia's within the test set that are correctly classified. Also if $W_{\text{false-Ia}}$ is set at one, then the second fraction represents the Ia purity – the proportion of test supernovae that we classify as Ia's that are actually Ia's. Kessler et al. [2010] suggested that $W_{\text{false-Ia}}$ be set at 3. This creates an extra penalty for incorrectly classifying a non-Ia as a Ia. With $W = 3$, the second fraction then represents the Ia pseudo-purity. We use this suggested value throughout to evaluate the performance of the three classifiers.

Since the proportion of Ia’s is ~70% within the training set and the Ia proportion is ~30% within the test set, the cross validation estimate of eqn 8 will be biased. We instead use a balanced classification rate to estimate the score. This entails using cross-validation to estimate the Ia efficiency and the non-Ia efficiency (denoted $\xi_{\text{Ia}}$ and $\xi_{\text{non-Ia}}$ respectively). Consequently, the cross-validation estimate using a balanced classification rate will be:

$$S_{\text{FoM-Ia}} = \xi_{\text{Ia}} \times \frac{L \times p_{\text{Ia}} \times \xi_{\text{Ia}}}{(L \times p_{\text{Ia}} \times \xi_{\text{Ia}}) + (W_{\text{false-Ia}} \times (L \times (1 - p_{\text{Ia}}) \times (1 - \xi_{\text{non-Ia}})))}$$

where $L$ is the total number of supernova in the test set and $p_{\text{Ia}}$ is the expected proportion of Ia's in the test set. In calculating the above score, we set $p_{\text{Ia}} = 0.3$. Given the above means of predicting the classifier performance, we now turn to the implementation of the three classifiers.

5.2.1 Nearest Neighbours (NN)

The Nearest Neighbours (NN) classifier simply compares the distances between the candidate supernovae and the pool of known Ia’s and non-Ia’s in the BAGIDIS coefficient space. The candidate is then classified in the same class as its closest matching supernova within the training set. The distance
between the i-th and j-th supernovae can easily be modified to account for the heteroskedasticity within the dataset. Let:

\[ d(d_i, d_j) = \sum_{k=1}^{D} \frac{|d_{i,k} - d_{j,k}|^2}{\sqrt{\sigma_k^{(i)^2} + \sigma_k^{(j)^2}}} \]

where \( d_i \) is the D-dimensional input vector that contain the relevant BAGIDIS coefficients for the i-th supernova and \( \sigma_k^{(i)^2} \) is the variance of the k-th BAGIDIS coefficient for the i-th supernova – as estimated from the Monte Carlo study described in sections 2.2 and 2.3. This distance measure can be calculated for all pairs of supernova that straddle the test and training sets. The only tuning parameter is the number of BAGIDIS coefficients \( D \) used to calculate the distance. This can be estimated using the balanced classification rates.

5.2.2 Support Vector Machines (SVMs)

Support Vector Machines (SVMs) are non-parametric classifiers that are known to be very competitive for many classification problems. They can also be relatively easily modified to account for the heteroskedasticity of the data. In addition, SVMs tend to be less susceptible to the curse of dimensionality than NN classifiers. We thus also applied SVM to the SNPC dataset. SVMs select a classification boundary that maximizes the margin between the boundary and the margin support vectors (selected observations with the dataset). For more details on SVMs see [Hastie et al. 2003]. SVMs allow for non-linear decision boundaries via the use of kernels. Kernels are a measure of similarity between two points within the input space. Amongst the most popular kernels is the Radial basis kernel. Here:

\[ K(d_i, d_j) = \exp \left( -\gamma \|d_i - d_j\|^2 \right) \]

where \( \gamma \) is a tuning parameter. The radial basis function implicitly assumes that the data is homoskedastic. We thus modify the kernel to account for the SNPC dataset’s heteroskedasticity.

\[ K(d_i, d_j) = \exp \left\{ -\gamma \sum_{k=1}^{D} \frac{(d_{i,k} - d_{j,k})^2}{\sqrt{\sigma_k^{(i)^2} + \sigma_k^{(j)^2}}} \right\} \]

where \( d_i = (d_{i,1}, ..., d_{i,D})^T \) is a D-dimensional input vector that contain the relevant BAGIDIS coefficients. Many SVM packages can accept pre-computed kernels as arguments, which makes it relatively straightforward to run an SVM using the above non-standard kernel.

When running the classifier, there are three tuning parameters whose values may be selected using the cross-validated balanced classification rates: \( D \) (the number of BAGIDIS coefficients to use as input within the SVM), \( \gamma \) (the kernel parameter) and \( C \) (a regularisation parameter that determines how tolerant the SVM classification boundary is to misclassification errors within the training set). The larger \( C \) is, the less tolerant we are of misclassification errors but the more likely we are to overfit the data. We run a grid search to choose the optimum values for \( \gamma \) and \( C \) on a logarithmic scale. We use cross-validation to predict the score at the parameter values: \( C = 2^{-5}, 2^{-3}, ..., 2^5 \) and \( \gamma = 2^{-15}, 2^{-13}, ..., 2^3 \). After running the grid search on a very large range of values, we can then narrow in on the sub-space where the classifier is performing best.

5.2.3 Ranked Probability Classifier

The ranked probability classifier is simple to implement. For each supernova within the test set, we calculate its probabilistic similarity to each of the training set supernova. The number of BAGIDIS coefficients to use for building the classifier (\( D \)) may be chosen using the cross-validated balanced classification rates. In addition, due to the false tag penalty within eqn [5] we may choose to only classify a supernovae with a BAGIDIS coefficient vector \( d_i^* \) as a Ia if:

\[ \log p(d_i|d_{IA}^*) > \log p(d_i|d_{non-IA}^*) + V \]

where \( d_{IA}^* \) and \( d_{non-IA}^* \) are the closest matching Ia and non-Ia supernovae respectively and \( V \) is a tuning parameter that can be chosen by cross-validation.

6 RESULTS FOR THE CLASSIFIERS

6.1 Results with the true, non-representative training set

We attempt to classify all test supernovae that have time spans greater than 100 days without using the redshift information. 54% of supernovae within the SNPCC test dataset satisfy this requirement. In current as well as future photometric surveys, we expect the vast majority of the supernovae captured to satisfy this requirement.

Table 1 shows the results for the NN classifier when training on various values of \( D \). We find that the optimum number of coefficients to train on is two. The two coefficient NN classifier also gave the best cross-validated score. Though only 57% of the Ia supernovae are identified, the objects that are classified as Ia’s have a very high purity (> 90%). Table 2 shows the results for the SVM classifier. The SVM tuning parameters \( \gamma \) and \( C \) were selected by cross-validation on the initial log scale before performing a finer grid search in the best performing region. This finer grid search was also done by cross-validation for the \( \gamma \) parameter but did not help in selecting the optimal \( C \) on a finer scale. The best performing score of 0.413 is misleadingly optimistic. If \( C \) is decreased or increased from 0.727 by even 0.001, the score drops to roughly ~ 0.300. The SVM classifier performs worse than the NN classifier. This is perhaps due to it’s sensitivity to the non-representativeness of the training set. Indeed, since for \( D = 3 \) the SVM’s classification performance is very sensitive to the value of \( C \), it appears that there is a high degree of variability in the SVM’s classification rule. In the next section, we compare the relative performances of the three classifiers on a representative training set.

Table 3 shows the performance of the Ranked Probability Classifier proposed in section 3. The Ranked Probability Classifier is the best performing classifier for all values of \( D \). One of the difficulties in implementing the classifier is that it provides perfect classification of the training set for a wide range of values of the tuning parameter \( V \). Table 4 shows its performance for both when \( V = 0 \) (i.e. we classify according to which class has the higher probability) as well as displaying the value of \( V \) for which the score on the test set is at its highest. The score increases when we make it easier to classify a point as a Ia – as can be seen in the fact
that the optimum $V$ for all values of $D$ is negative. Reassuringly however, the score does not drop by much if we just use $V = 0$ as our default.

6.2 Results with a representative training set

We now turn our attention to the performance of the classifiers if we were able to run the classifiers on a representative training set. Since it is unrealistic to have a truly representative training set, we wish to discern the robustness of the three classifiers to the non-representativity of the training set. Also, if classification algorithms are egregiously affected by the non-representativity, it would motivate the investment of substantial resources towards the production of a more representative training set.

Of the roughly 10,000 supernovae with a timespan of at least 100 days, less than 5% have known class. The organisers of the challenge later released the classes of all supernovae, so we can randomly assign roughly 5% of supernovae to have a known class and the remaining 95% to be part of the test set. Subsequently we train all three classifiers on this new training set. Recall that we previously avoided using the first coefficient in all three classifiers. As expected, the performance of all three classifiers improves when using a representative training set, including the information within the first coefficient would be overly sensitive to the training supernovae being brighter on average than the test supernovae. Now that we are training on a representative training set, including the information within the first coefficient will improve performance. However in the interest of being able to directly compare the results with the classifiers that use the true training set, we still choose to avoid using the first coefficient in all three classifiers.

Tables 3, 4 and 5 show the the performances on a representative training set for the NN, SVM and Ranked Probability classifiers respectively.

| Table 1. | Classification results for the Nearest Neighbours (NN) classifier for different number of $BAGIDIS$ coefficients. The highest score is shown in bold. |
|----------|----------------------------------------------------------------------------------------------------------------------------------|
| $D$ | Efficiency | Purity | Ps. Purity | Score |
| 2 | 57.4% | 91.2% | 77.6% | 0.445 |
| 3 | 53.1% | 92.2% | 79.8% | 0.424 |
| 4 | 57.4% | 87.1% | 69.3% | 0.398 |
| 5 | 58.8% | 86.3% | 67.8% | 0.398 |

| Table 2. | Classification results for the Support Vector Machine (SVM) classifier for different number of $BAGIDIS$ coefficients. The corresponding $\gamma$ and $C$ have been selected at the log-scale using the cross-validated balanced classification rates. The highest score is shown in bold. |
|----------|----------------------------------------------------------------------------------------------------------------------------------|
| $D$ | $\gamma$ | $C$ | Efficiency | Purity | Ps. Purity | Score |
| 2 | 0.009 | 0.5 | 75.3% | 64.0% | 37.2% | 0.280 |
| 2 | 0.009 | 0.483 | 85.7% | 61.8% | 35.0% | 0.300 |
| 3 | 0.010 | 0.036 | 59.6% | 60.2% | 33.5% | 0.200 |
| 3 | **0.010** | **0.727** | **68.4%** | **82.0%** | **60.3%** | **0.413** |
| 4 | 0.018 | 0.043 | 61.5% | 67.4% | 40.8% | 0.238 |
| 4 | 0.014 | 1.006 | 69.9% | 69.2% | 42.8% | 0.269 |
| 5 | 0.014 | 0.910 | 71.2% | 68.0% | 41.4% | 0.295 |

| Table 3. | Classification results for the Ranked Probability classifier for different number of $BAGIDIS$ coefficients. $V$ is a tuning parameter (see section 5.2.3). The highest score is shown in bold. |
|----------|----------------------------------------------------------------------------------------------------------------------------------|
| $D$ | $V$ | Efficiency | Purity | Ps. Purity | Score |
| 2 | 0 | 72.6% | 84.5% | 64.4% | 0.468 |
| 2 | -1.1 | 75.0% | 83.7% | 63.1% | 0.473 |
| 3 | 0 | 75.2% | 83.6% | 63.0% | 0.474 |
| 3 | -2.3 | 79.7% | 81.7% | 59.8% | 0.477 |
| 4 | 0 | 75.7% | 83.0% | 62.0% | 0.470 |
| 4 | -1.3 | 78.6% | 82.6% | 61.2% | 0.481 |
| 5 | 0 | 77.5% | 83.1% | 62.1% | 0.481 |
| 5 | **-1.4** | **80.5%** | **82.4%** | **60.9%** | **0.490** |

| Table 4. | Classification results for the Nearest Neighbours (NN) classifier on a representative training set for different number of $BAGIDIS$ coefficients. The highest score is shown in bold. |
|----------|----------------------------------------------------------------------------------------------------------------------------------|
| $D$ | Efficiency | Purity | Ps. Purity | Score |
| 2 | 70.7% | 84.2% | 64.0% | 0.452 |
| 3 | 76.7% | 84.9% | 65.2% | 0.500 |
| 4 | **76.4%** | **86.2%** | **67.6%** | **0.517** |
| 5 | 78.1% | 85.3% | 65.8% | 0.514 |

| Table 5. | Classification results for the Support Vector Machine (SVM) classifier on a representative training set for different number of $BAGIDIS$ coefficients. The corresponding $\gamma$ and $C$ have been selected at the log-scale using the cross-validated balanced classification rates. The highest score is shown in bold. |
|----------|----------------------------------------------------------------------------------------------------------------------------------|
| $D$ | $\gamma$ | $C$ | Efficiency | Purity | Ps. Purity | Score |
| 2 | 0.006 | 0.031 | 71.5% | 80.7% | 58.3% | 0.417 |
| 2 | 0.006 | 0.043 | 76.6% | 78.7% | 55.1% | 0.423 |
| 3 | 0.016 | 0.125 | 77.2% | 73.8% | 48.5% | 0.374 |
| 3 | 0.016 | 0.109 | 78.1% | 75.3% | 50.5% | 0.394 |
| 4 | 0.015 | 0.125 | 77.5% | 73.1% | 47.6% | 0.369 |
| 4 | 0.015 | 0.106 | 78.6% | 75.7% | 50.9% | 0.400 |
| 5 | 0.005 | 0.031 | 71.4% | 81.2% | 59.0% | 0.421 |
| 5 | **0.005** | **0.036** | **74.9%** | **80.2%** | **57.4%** | **0.430** |

Note: The tables show the performances on a representative training set for the NN, SVM and Ranked Probability classifiers respectively. As expected, the performance of all three classifiers improves when using a representative training set. However, even with a representative training set, the SVM still is the worst performing classifier and the Ranked Probability classifier is still the top performing classifier. We also wish to compare the relative robustness of the three classifiers to the non-representativity of the training set. We thus look at the percentage increase in the score of each classifier when using a representative dataset. Note for the SVM classifier, we are able to reliably choose the value of the tuning parameter $C$ at the log-scale but not on a finer scale. Hence in calculating the percentage increase for the
SVM we only consider the scores where the percentage increase for the cases where choosing 2 of the resulting classification boundary often increases exponentially with the dimensionality of the space. High dimensional spaces are problematic as the algorithmic object will often exist in a very high dimensional area. We believe that classifiers that base the classification rule for a candidate object on the closest matching training object in each of the classes will tend to be more robust to the non-representativity of the input space since training data that are the closest matches to the test set should be more representative of the test set. The nearest neighbours classifier (NN) is an example of such a classifier. We also propose the Ranked Probability classifier. This classifier uses a probabilistic measure of similarity to determine the closest matching training object in each class.

We applied the proposed methodology to the Supernova Photometric Classification Challenge (SNPCC). To the best of our knowledge, this is the first analysis that does not require the curves to be re-aligned so that peak brightness is at time zero. Such a re-alignment can be computationally expensive and makes assumptions about the form of the supernova light curves. The challenge consisted of trying to identify the Ia supernova based on a series of flux measurements. The quality of this functional data varied drastically between the supernovae; both in terms of the number of flux measurements and the measurement error bars. Despite this, we were able to standardize the dataset by fitting a spline to the data and taking the resulting spline estimates on an equidistant grid. The data within this series of n spline estimates was compressed into $D$ BAGIDIS coefficients ($D \ll n$). A t-SNE plot revealed that the second to the sixth BAGIDIS coefficients provided excellent discriminatory information on the class of the supernova. We then applied an NN, an SVM and the Ranked Probability classifier to all those supernovae with a timespan greater than 100 days. This constituted 54% of the SNPCC test dataset and we expect the percentage to be much larger in future photometric surveys. Both the NN and the Ranked Probability classifier provided highly competitive results with the Ranked Probability classifier offering over 80% Ia efficiency and purity. We also noted that both classifiers appeared to be more robust to the non-representativity of the training set than the SVM classifier. We speculate that this could be due to both classifiers basing the classification rule on a subset of the training set that is more representative of the test set. Though the methodology presented in this paper was applied to light curve data, it can also be readily applied to other types of astronomical functional data such as spectra. The BAGIDIS methodology could potentially also be used to summarise the light curves captured by future large synoptic sky surveys.

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