**SNIascore: Deep-learning Classification of Low-resolution Supernova Spectra**

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**Abstract**

We present SNIascore, a deep-learning-based method for spectroscopic classification of thermonuclear supernovae (SNe Ia) based on very low-resolution ($R \sim 100$) data. The goal of SNIascore is the fully automated classification of SNe Ia with a very low false-positive rate (FPR) so that human intervention can be greatly reduced in large-scale SN classification efforts, such as that undertaken by the public Zwicky Transient Facility (ZTF) Bright Transient Survey (BTS). We utilize a recurrent neural network architecture with a combination of bidirectional long short-term memory and gated recurrent unit layers. SNIascore achieves a <0.6% FPR while classifying up to 90% of the low-resolution SN Ia spectra obtained by the BTS. SNIascore simultaneously performs binary classification and predicts the redshifts of secure SNe Ia via regression (with a typical uncertainty of $<0.005$ in the range from $z = 0.01$ to $z = 0.12$). For the magnitude-limited ZTF BTS survey ($\sim$70% SNe Ia), deploying SNIascore reduces the amount of spectra in need of human classification or confirmation by $\sim$60%. Furthermore, SNIascore allows SN Ia classifications to be automatically announced in real-time to the public immediately following a finished observation during the night.

**Unified Astronomy Thesaurus concepts:** Supernovae (1668); Type Ia supernovae (1728); Neural networks (1933); Classification (1907); Surveys (1671)

1. **Introduction**

Modern time-domain surveys, such as the Zwicky Transient Facility (ZTF), Bellm et al. 2019a, 2019b; Graham et al. 2019; Masci et al. 2019; Dekany et al. 2020, the All-Sky Automated Survey for Supernovae (ASAS-SN; Shappee et al. 2014) and the Asteroid Terrestrial-Last-Alert System (ATLAS; Tonry et al. 2018b), are now finding tens of thousands of transients every year.

However, without spectroscopic classifications, these discoveries are of limited value (Kulkarni 2020). The ZTF Bright Transient Survey (BTS; Fremling et al. 2020; Perley et al. 2020) is addressing this through the deployment of a fully automated very-low-resolution spectrograph, the Spectral Energy Distribution Machine (SEDM; Blagorodnova et al. 2018; Rigault et al. 2019) mounted on the Palomar 60 inch telescope. SEDM is capable of obtaining spectra of several thousands of transients per year in the magnitude range between 18 and 19 mag. Currently, the goal of the BTS is to maintain spectroscopic classification completeness for all extragalactic transients detected by the public ZTF survey that become brighter than 18.5 mag ($\sim$1000 SNe per year; Perley et al. 2020).

The classifications from the BTS are made public on a daily basis via the Transient Name Server (TNS1). These classifications have up until now been based on manual matching of observed spectra to spectral templates using mainly the SuperNova IDentification (SNID; Blondin & Tonry 2007) code, along with careful inspection of each obtained spectrum. This makes classification of thousands of SNe a very time-consuming endeavor.

Due to their inherent brightness, the majority of the extragalactic transients discovered by a magnitude-limited survey will be thermonuclear supernovae (SNe Ia)12. Here we present SNIascore, a deep-learning-based method optimized to identify SNe Ia using SEDM spectra and determine their redshifts without any human interaction. The intended use case for SNIascore is to provide live spectroscopic classification of SNe Ia during the night when SEDM is observing for the BTS.

SNIascore is based on a recurrent neural network (RNN) architecture (see Sherstinsky 2020 for a review) with a combination of bidirectional long short-term memory (BiLSTM) and gated recurrent unit (GRU) layers. SNIascore is able to classify >80% of the SN Ia spectra that are observed by SEDM for the ZTF BTS with a false-positive rate (FPR) of <1%.

SNIascore was trained on SEDM data obtained by the ZTF BTS between 2018 and 2020 March, and validated and optimized with the BTS data set published by Fremling et al. (2020). A final

[11] https://wis-tns.org
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**Table 1**

| Class            | All | Training | Validation | Testing |
|------------------|-----|----------|------------|---------|
| SNIIa            | 2619 | 1526     | 607        | 486     |
| NotSNIIa         | 2931 | 1997     | 409        | 525     |
| H-rich CC SN     | 1285 | 751      | 312        | 222     |
| H-poor CC SN     | 585  | 393      | 94         | 98      |
| TDE              | 37   | 35       | 0          | 2       |
| CV               | 325  | 284      | 0          | 41      |
| Other            | 699  | 534      | 3          | 162     |

Note. All numbers refer to the number of unique spectra in each class. The “Other” class includes any spectra that do not fit the other classes, including galaxy spectra, active galactic nucleus (AGN) spectra, and spectra that BTS has been unable to classify as belonging to any known transient class.

test run is performed using data obtained between 2020 April and 2020 August, neither of which were part of the training or validation data sets. Our data sets are described in detail in Section 2. The network architecture is described in Section 3, and the training and optimization procedure used for SNIIascore is described in Section 4. The performance is evaluated and compared to SNID and the previously published deep-learning method DASH (Muthukrishna et al. 2019b) in Section 5. A discussion on the most likely false positives can be found in Section 6. The implementation of SNIIascore as part of the ZTF BTS is described in Section 7. Future development possibilities are also discussed in this section.

### 2. Data Sets

Our data consist of SEDM spectra of transients detected by ZTF, which were followed up and classified by the BTS (Fremling et al. 2020; Perley et al. 2020) using the GROWTH Marshal (Kasliwal et al. 2019). These spectra are of low resolution (R ~ 100) and cover a typical wavelength range of 3800–9150 Å within 209 wavelength bins (see Section 2.2). Although the BTS focuses on extragalactic transients, some spectra eventually turn out to be of Galactic sources (e.g., cataclysmic variables; CVs). We include both spectra of Galactic and extragalactic transients in our data sets. The data set we use here contains 5550 SEDM spectra of 3463 individual transients obtained between 2018 March and 2020 August. A breakdown of the various classes of transients included in our data set can be found in Table 1. We split our full data set into three parts: training data, validation and optimization data, and final performance testing data.

For the purpose of training SNIIascore (Section 4) we group the data into two classes: real SNIIa (SNIIa; 1526 spectra of 1090 SNIIa) and everything else (NotSNIIa; 1997 spectra of 1121 transients). The training data were collected between 2018 March 7 and 2020 March 1, but we exclude any data that are part of our validation set, which was also collected during 2018.

To validate and optimize SNIIascore (Section 4) we use 1016 spectra of 648 SNII, which are part of the BTS sample from 2018 published in Fremling et al. (2020; hereafter the BTS18 sample). The BTS18 sample was chosen for validation because the classifications and redshifts of the SNII in this sample have been carefully vetted by humans.

For final performance testing we use all BTS spectra collected between 2020 March 2 and 2020 August 5 (1011 spectra of 632 transients). The classifications in this testing sample are sufficiently accurate to evaluate the performance of SNIIascore, but they are subject to minor future changes as work on the BTS proceeds.13

For training SNIIascore for redshift prediction, we only use spectra of SNIIa and also impose a cut on the quality of the known redshifts. We only use SNIIa where the redshift is known to three decimal places or more.14 Both redshifts derived from broad SN features and host-galaxy emission lines are used. The training set for redshift prediction consists of 891 spectra of 630 SNIIa, which is increased to 12,810 spectra through data augmentation (Section 2.1).

The properties of the training, validation, and testing data sets are illustrated in Figure 1. The redshift range covered by our full data set is $z = 0.008$ to $z = 0.126$. However, 98% of our data fall within the range $z = 0.01$ to $z = 0.11$. The phases of our spectra with respect to maximum light determined from the lightcurves of our transients fall within $−20$ days to $+30$ days, with 97% of the data within $−20$ days to $+20$ days. Thus, the range we can expect SNIIascore to perform reliably within is $z = 0.01$ to $z = 0.11$ for spectra obtained within $±20$ days of maximum brightness. We use the time of maximum light from the BTS Sample Explorer (Perley et al. 2020), which records the brightest actual measurement in the lightcurve of each transient. This measurement can be in the $g$ or $r$ band, depending on which band is brighter.

### 2.1. Training Set Augmentation

For the classification component of SNIIascore we can achieve excellent performance without any need for data augmentation (Section 5). We also do not need to de-redshift our spectra as part of the preprocessing procedure that is performed before training or when classifying new data (Section 2.2). However, in order to use regression to predict the redshifts of SNIIa identified by SNIIascore without

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13 The live BTS sample, which is updated daily, can be accessed through the BTS Sample Explorer at https://sites.astro.caltech.edu/ztf/bts/explorer.php.

14 This typically means that a spectrum of the host galaxy exists in the literature (e.g., on NED), the SEDM spectrum was of very high signal-to-noise ratio (S/N) with very good SNID template matches or clear host-galaxy lines, or that we have obtained a spectrum of the SN from a telescope that offers a higher resolution compared to SEDM.
introducing systematic bias, a weighting scheme, or data augmentation, is needed. This is due to the shape of the redshift distribution of our data set (Figure 1); there are very few spectra for SNe Ia at both $z < 0.03$ and $z > 0.09$.

We have found that a simple augmentation procedure, which equalizes the redshift distribution of the training sample, can offer excellent redshift prediction performance with negligible bias. We perform the following steps: for each redshift bin of size 0.001 from zero to 0.13 we search for spectra in our unaugmented data set with similar redshifts (within $z \pm 0.005$ to $z + 0.01$). We then randomly pick spectra from the matches and apply new redshifts by shifting the wavelengths of the original spectra to randomly selected redshifts within $\pm 0.001$ of the redshift bin center (the spectra are not de-redshifted to the rest frame as part of this process to reduce loss of information at the edges of the spectra). We repeat this process until we have roughly 100 spectra in each redshift bin, for a total of 12810 spectra. To reduce overfitting and lessen the impact of repeating a small number of spectra at the edges of the redshift distribution, we introduce noise from a normal distribution to each added spectrum. The final redshift distribution for our training data used for redshift regression is shown in Figure 1.

### 2.2. Preprocessing of Spectra

Due to small variations in the final wavelength coverage of the SEDM spectra included in our data set, we interpolate all spectra to a uniform wavelength grid, which spans the largest possible wavelength range without a need for extrapolation for any individual spectrum. This range is 3800–9150 Å with 209 wavelength bins of size 25.6 Å.

After interpolation we normalize each spectrum by division with its median value. The normalized spectrum is then divided by its continuum (a heavily smoothed version of the normalized spectrum computed with robust local polynomial regression using weighted linear least squares and a second-degree polynomial model). Finally, to center the preprocessed spectra around zero, we subtract a constant value of one from each normalized and continuum divided spectrum. This process is illustrated in Figure 2.

We have found that this preprocessing procedure significantly outperforms a simple division by the median, directly dividing by a smoothed continuum, or dividing by the median and then subtracting the smoothed continuum. We have also investigated the effect of suppressing the bluest and reddest parts of the spectra (which can be very noisy in data from many instruments), as in Muthukrishna et al. (2019b). This decreases the overall performance and is not needed for SEDM data.

Effectively our preprocessing procedure flattens the spectra (removes any temperature gradient or host-galaxy continuum emission contribution), makes emission lines positive and absorption lines negative, and normalizes the strengths of the features.

### 3. Neural Network Architecture

For our final SNiascore network that we have arrived at after following the optimization procedure described in Section 4, we have used an RNN architecture (see e.g., Sherstinsky 2020) consisting of a combination of BiLSTM and GRU layers, with 32 hidden units in each layer (Figure 3). We employ significant dropout both during training and prediction. We use a dropout of 40% immediately following the input layer. This is followed by a BiLSTM layer and 45% dropout. After this we use a GRU layer and 35% dropout and then another BiLSTM layer and 25% dropout. This last BiLSTM layer is followed by a fully connected layer with two outputs, a softmax layer and a classification layer that computes the cross-entropy loss for the SNIa and NotSNIa classes.

For redshift regression the softmax and classification layers are replaced by a custom regression layer, which uses the root of the mean bias error (ME; Equation A4) as the backward loss function and the derivative of the mean absolute error (MAE; Equation A4) as the forward loss function. We have found that this custom regression layer gives the least systematic bias and best overall performance among the loss functions we have tested (Section 4).

To estimate uncertainties when predicting both the SNiascore ($\sigma_{\text{SNiascore}}$) and redshift ($\sigma_z$), we use a simple Monte Carlo (MC) method. Because dropout is used during prediction, the results of subsequent predictions based on the same input spectrum will be different. We rerun each prediction 100 times and adopt the standard deviation of the results as the uncertainty of the prediction (Gal & Ghahramani 2015).

### 4. Training and Optimization

#### 4.1. Classification Optimization

We have implemented the architecture for SNiascore described in Section 3 using the MATLAB Deep Learning Toolbox™. For training we use the adaptive learning rate optimization algorithm Adam (Kingma & Ba 2014) with the following settings:  

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15 We add between zero and one standard deviation of noise. The standard deviation is estimated as the standard deviation of each spectrum after subtraction of the broad features using a heavily smoothed spectrum.

16 Lower weights are assigned to outliers in the regression; outliers at $>6\sigma$ are assigned zero weight.
Figure 3. Network architecture of SNIascore. We utilize heavy dropout throughout the network and a combination of two BiLSTM layers surrounding one GRU layer. For regression the final softmax and classification layers are replaced by a regression layer.

1. MiniBatchSize: 256
2. InitialLearnRate: 0.005
3. LearnRateSchedule: none
4. L2Regularization: $10^{-4}$
5. GradientThreshold: 2
6. Shuffle: every-epoch
7. SequenceLength: longest.

For validation and optimization we use the BTS18 data set. To arrive at the architecture described in Section 3 and the optimal hyperparameters listed above, we have followed this procedure:

1. We construct a grid of networks with an initial dropout layer following the input layer and then one to five BiLSTM layers with dropout layers between each layer.
2. For each network we create a grid of subnetworks with a range of hyperparameters, dropout from 0.15 to 0.45 for each dropout layer, and hidden units from 8 to 128 for each BiLSTM layer.
3. We explore learning rates in the range 0.001–0.01 and mini batch sizes of $2^n$ in the range 16–512 for each network.

This optimization procedure showed that a constant learning rate of 0.005 and a mini batch size of 256 seems to perform universally well on our data, regardless of the other hyperparameters. Changes to the learning rate (including introducing adaptive learning rate schemes), gradient threshold, and L2 regularization listed above have negligible impact on the final performance. As long as the network retains the ability to converge during training, it remains possible to pick an optimal epoch from the training sequence where the performance is very similar.

We do not try to optimize for the required physical training time or number of epochs required to achieve a good performance. We do not employ any stopping condition based on the loss. Instead, each network is trained past the point of overfitting (at least 325 epochs), and we save the state of each network at every training epoch. We then choose and compare the optimal epoch (typically found between epochs 200 and 300) for every individual network based on the performance on the BTS18 validation set, according to a combination of two metrics: (i) the total number of successful classifications for an FPR of 0%, and (ii) the total number of successful classifications for an FPR of 1%. To determine the optimal FPR and true-positive rate (TPR) of an individual network, all possible cuts on SNIascore and $\sigma_{\text{SNIascore}}$ are evaluated. These metrics direct the optimization procedure toward hyperparameters and a network structure that results in a low FPR, with a high TPR being secondary. We consider FPR < 1% to be a hard requirement for autonomous spectroscopic classification. It also allows all the trained networks to be quantitatively compared based on their optimal epochs and SNIascore cuts.

The architecture that resulted from this optimization procedure was 3 BiLSTM layers with 32 hidden units in each layer, in combination with heavy dropout (0.40, 0.45, 0.35, and 0.25 for the 4 dropout layers from first to last). As a final step, we investigated the effect of replacing the BiLSTM layers with GRU layers in all possible combinations and eventually arrived at our final SNIascore network described in Section 3, where the middle BiLSTM layer is replaced by a GRU layer. The effect of this is a reduced number of false positives with high SNIascore values.

For our optimized classification network, Figure 4 shows the SNIascore distribution for known SNe Ia and other transients in the BTS18 validation sample. We find that 90% of the SNe Ia in this sample are identified by SNIascore with a 0.6% FPR. The performance of our final SNIascore network is investigated in detail and tested on independent data in Section 5.

4.2. Redshift Regression Optimization

For redshift regression we use the same optimal SNIascore architecture found above with the softmax and classification layers replaced with a regression layer (Figure 3). However, we have found that the standard deep-learning mean square error (MSE; Equation (A1)) or MAE loss functions result in significant redshift-dependent systematic bias, even after data augmentation to equalize the redshift distribution of the training data (Section 2.1). Because of this, we perform an additional optimization step for the redshift regression network, which consists of an investigation of various forward and backward loss functions in the final regression layer of the network.

The forward loss functions that we have investigated are the MAE, the mean percentage error (MPE; Equation (A3)), the MSE, the MBE, and all possible combinations of these in pairs, with both the derivative of the MAE and MSE as the backward loss.

The MPE, MBE, and the root square of these can all give good results. Differences are minor, but we have selected the root square bias error ($\sqrt{\text{MBE}^2}$) as the forward loss for our final network and the derivative of the MAE as the backward loss, based on optimizing (i) the percentage of SNe with redshift residuals less than 0.005 and 0.01 for the BTS18 validation sample, and (ii) how close the predictions on the full validation set adhere to a linear relationship with respect to the known redshifts ($y = ax + b$ where $a$ should be as close to 1 as possible and $b$ as close to zero as possible). Each network was trained past overfitting (at least 525 epochs) and the optimal epoch (typically found between epochs 400 and 500) was selected after the fact for comparison among the networks.

5. Performance

In order to evaluate the classification performance of SNIascore we have performed comparisons to SNID (Blondin & Tonry 2007) and DASH (Muthukrishna et al. 2019b) (Section 5.1). We also evaluate the accuracy of the
associated redshift predictions for the SNe Ia that are identified by SNIascore by comparing to host-galaxy redshifts and redshifts determined after manual inspection with SNID (Section 5.2).

5.1. Classifications

Our goal with SNIascore is a low-enough FPR (<1%) so that the classifications do not require any human confirmation. To achieve this, using the standard value of SNIascore > 0.5 to classify a spectrum as that of an SN Ia is not adequate. Instead, we evaluate all possible cuts on SNIascore and σSNIascore to find a combination of cuts that result in the desired performance.

Receiver operating characteristic (ROC) curves for SNIascore (computed by varying SNIascore and σSNIascore), SNID (by varying the rlap parameter threshold), and DASH (by varying rlap and the softmax score threshold) are shown in Figure 4. Based on this comparison it is clear that SNIascore significantly outperforms these alternatives. Between 83% (testing data) and 90% (validation data) of the SN Ia spectra that SEDM observes for the ZTF BTS can be classified with FPR ≈ 0.5%. This performance is achieved for the cuts SNIascore > 0.6 and σSNIascore < 0.3, which produce F1-scores of 0.95 and 0.91 for the validation and testing data sets, respectively.

SNID can achieve a very low FPR for SN Ia spectra when rlap > 10. However, only ≈50% of the spectra could potentially be automatically classified (FPR < 1%) without supervision with SNID. The deep-learning-based DASH only reaches a low-enough FPR to enable automatic classification when applied to SEDM spectra for 24% of spectra for any cuts on rlap and the softmax score. This is likely due to the fact that no spectra with similarly low resolution to those produced by SEDM were part of the DASH training data. More conservative cuts at SNIascore > 0.8 and σSNIascore < 0.275 result in zero false positives on the validation and training data, with F1-scores of 0.91 and 0.86 for the validation and testing data sets, respectively. Depending on the use case, either choice of these cuts may be suitable. For live reporting of classifications to the community via TNS, the stricter cuts seem

\[ \frac{N_{\text{true}}}{N_{\text{tot}}} \]

\[ \text{FPR} \]
appropriate (Section 7). We summarize our recommended SNIascore cuts and the performance compared to SNID and DASH in Table 2. An investigation of how the classification performance of SNIascore varies depending on data quality can be found in Appendix C.

Finally, we note that a low SNIascore does not necessarily mean that the transient being observed cannot be an SN Ia. There are multiple reasons for low scores, for example, low S/N, strong galaxy light contamination, and significant contamination from cosmic rays that failed to be removed during data reduction. As such, a score that is lower than the cuts discussed above should generally be interpreted as that the spectrum cannot be used to make a confident SN Ia classification, and nothing further. Only when the SNIascore is extremely low (<0.01), is there significant statistical power to support a conclusion that the transient associated with the respective spectrum cannot be an SN Ia. We have made no attempt at optimizing the FNR for low scores, but based on the FNR curves for the validation and testing data sets (Figure 4), SNIascore < 0.01 corresponds to FNR ≈ 1%.

5.2. Redshifts

Because only SN Ia spectra were used during training for redshift prediction (Section 4.2), we only perform predictions for spectra that pass the stricter cuts that optimize SNIascore for low FPR (Table 2). Any false positives will receive inaccurate redshift predictions, so they need to be avoided.

To evaluate the accuracy of the redshifts predicted by SNIascore, we compare predictions for the BTS18 validation and testing data sets primarily to redshifts derived from host-galaxy spectra, and secondarily to redshifts determined from broad SN features through manual template matching using SNID which were performed for the BTS18 SEDM data in Fremling et al. (2020). The performance of the SNIascore redshift regression network is illustrated in Figure 5.

We find that the performance of our network optimized on the BTS18 validation data carries over very well to the testing data set; we find no evidence for systematic bias in the predictions from SNIascore for either data set. The absolute value of the difference between each prediction compared to the redshift from the host-galaxy spectrum is <0.005 for 77% of the spectra in the validation set and for 85% of the spectra in the testing data set. For 96% (validation data) and 100% (testing data), this difference is <0.01.

We find the mean of the redshift residuals, Δz = −0.0008, and the standard deviation, σz = 0.0046, for the BTS18 validation data when SNIascore is compared to redshifts from host-galaxy spectra. For the testing data we find Δz = 0.0006 and σz = 0.0035, when compared to redshifts from host-galaxy spectra. When SNIascore is compared to redshifts derived manually from broad SN features using SNID we find Δz = −0.0002 and σz = 0.0049 (only possible for the validation data). It has been previously found that σz = 0.005 when manual usage of SNID is compared to redshifts from host-galaxy spectra on the BTS18 sample (Fremling et al. 2020), which is consistent with what was found for higher-resolution spectra by Blondin & Tonry (2007). As such, we conclude that the performance of SNIascore is consistent with what can be done with SNID when each result and spectrum is manually inspected. A detailed investigation of the accuracy of automatic SNID SN Ia redshifts from SEDM is being performed for data up until 2021 (M. Rigault et al. 2021, in preparation).

6. Discussion

A few spectra in both our validation and testing data sets end up as false positives with the optimal TPR cut we suggest to be used with SNIascore (Table 2). Most of these spectra are of low quality or affected by strong host contamination (and uninteresting). However, there are also a few high-quality spectra. It is interesting to investigate these and the associated SNe in some detail.

In the testing set, ZTF20aatxryt (SN 2020eyj; Tonry et al. 2020) stands out. Our first SEDM spectrum, taken around maximum light of the SN on 2020 April 2, is of high quality and looks consistent with an SN Ia spectrum and gets an SNIascore of 0.7 ± 0.3. However, it later turned out that the ejecta from this SN crashed into circumstellar material, resulting in a flattening of its lightcurve and changing its spectrum into that of an SN Ibl (E. Kool et al. 2021, in preparation). The best matches in SNID are also SNe Ila with rlap > 10. However, by manually going through all the matches produced by SNID at rlap > 5, some SN Ic matches can also be found. Given the lightcurve evolution and the fact that SN Ia lightcurve models do not match well even around maximum light before the interaction sets in, it may be more likely that this was a stripped-envelope SN. We note that the ZTF BTS officially classified this as an SN Ia as part of our routine operations based on the SNID match to the peak spectrum (Dahiwale & Fremling 2020). Only later did we realize that the object was unusual.

In the validation set, ZTF18aaxmhvk (SN 2018cne; Tonry et al. 2018a) stands out (SNIascore = 0.71 ± 0.17). Our SEDM spectrum of this source, taken around maximum light on 2018 June 14, is of high quality and is also best matched to SNe Ia in SNID with very high rlap values. However, the lightcurve of ZTF18aaxmhvk is not consistent with that of a normal SN Ia. Based on this, and the fact that matches to SNe Ic can also be found in SNID, the BTS officially classified this as an SN Ic (Fremling & Sharma 2018). However, it could also have been a peculiar SN Ia. The SEDM spectrum is the only spectrum available for this source; we lack higher-resolution spectra and do not have any late-time spectra, which could have provided a conclusive answer.

In conclusion, the main contaminants that may produce high SNIascore values appear to be stripped-envelope SNe, and in particular events with SN Ic-like spectra. The BTS has been classifying roughly 30 SNe Ic per year (Perley et al. 2020).

| Method | TPR | FPR | Cuts* |
|--------|-----|-----|-------|
| SNIascore TPR | 0.90 | 0.006 | score > 0.6, σ < 0.3 |
| SNIascore FPR | 0.86 | 0.000 | score > 0.8, σ < 0.275 |
| SNID | 0.53 | 0.009 | rlap > 11.1 |
| DASH | 0.24 | 0.009 | rlap > 7.1, score > 0.5 |

Notes. For SNIascore TPR, SNID, and DASH we show the optimal TPR that can be had while maintaining FPR < 1%. For SNIascore FPR, we show the optimal TPR that can be had for FPR = 0.

*score refers to the softmax score for both SNIascore and DASH. For SNIascore, σ refers to σSNIascore.
Thus, as there is only one SN Ic spectrum that gives a high SNIAscore value in each of the validation and testing data sets, it appears that only in rare cases do SNe Ic cause confusion for the BTS (which probes \( z < 0.1 \) for core-collapse SNe). Furthermore, if the lightcurves of these SNe are taken into account it appears relatively straightforward to remove such objects from any sample of SNe Ia that is required to be devoid of both core-collapse SNe and highly peculiar SNe Ia.\(^{19}\)

For future versions of SNIAscore, we plan to add the option to include lightcurve information as input. Looking at objects that change SNIAscore significantly when the lightcurve is included or excluded may turn out to be a way to identify rare events that would warrant further follow-up from larger facilities. However, the intended use case of SNIAscore is primarily to provide live classifications when SEDM is observing. As such, a potential limitation is the fact that we do not necessarily have much lightcurve information available at the time of the spectral observation by SEDM. A significant fraction of our spectra are observed before maximum light (Figure 1), and full lightcurves cannot be leveraged.

Some work on machine-learning methods for early photometric classification has been done (e.g., RAPID; Muthukrishna et al. 2019a). However, we have not been able to reproduce the expected performance in Muthukrishna et al. (2019a) on live ZTF data. Work on lightcurve classifiers trained on ZTF data is in progress, which we plan to eventually combine with SNIAscore and the redshift predictions that we already produce. Work on a deep-learning classifier capable of distinguishing between more SN subtypes while maintaining a comparably low FPR is also ongoing (Y. Sharma et al. 2021, in preparation).

7. Summary and Implementation

With SNIAscore we have shown that a simple RNN architecture (Section 3) can be used to create a method that is capable of fully automatic classifications of low-resolution SN Ia spectra obtained by the SEDM instrument. Most importantly, SNIAscore achieves a consistently low FPR of \(<1\%\) when classifying up to 90\% of the SN Ia spectra that SEDM observes during routine operation (Section 5.1). For securely classified SN Ia spectra, SNIAscore also achieves excellent redshift regression performance, comparable to determining redshifts by carefully running each spectrum manually through SNID. Typically, the redshift predictions made by SNIAscore differ by less than \( z \pm 0.005 \) when compared to host-galaxy redshifts (Section 5.2).

Starting from 2021 April 15 we have fully deployed SNIAscore as part of the automated ZTF SEDM pipeline (Fremling et al. 2021). This includes automatic reporting of confident SN Ia classifications and redshift estimates to TNS. With this, we have achieved automation all the way from (spectroscopic) observation to data reduction and spectroscopic classification for the first time.

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\(^{19}\) We note that with the low resolution of SEDM, it is not possible to spectroscopically separate between the various SN Ia subtypes, except in the most clear-cut cases (see Fremling et al. 2020).
The first fully automated SEDM SN Ia classification was sent for SN 2021jib (SNIascore 2021). This report was sent within roughly 10 minutes after the finished SEDM exposure. With automated ZTF candidate vetting and automatic triggering of SEDM, which is currently possible through AMPLE (Nordin et al. 2019) and the Fritz marshals (Duev et al. 2019; van der Walt et al. 2019), a fully automated imaging (ZTF) and spectroscopic (SEDM) survey is now becoming realistic.

Initially for our TNS reports, we are using a very conservative SNIascore threshold of 0.9. Based on the performance evaluation in Section 5.1, we plan to eventually relax this to a cut on the SNIascore of >0.8 combined with a cut on \( \sigma_{\text{SNIascore}} \) of <0.275. This will eliminate the need for human interaction in the classification process for the majority of BTS SNe Ia (\( \approx 80\% \) based on our testing data or \( \approx 90\% \) based on our validation data), which translates to >50\% of BTS SNe.

We do not find significant variations in the FPR as a function of time or overall data quality (Appendix C), based on the data currently available to us. However, we note that the corresponding TPR (e.g., Table 2) and the numbers above are subject to changing observing conditions and the condition of the primary mirror of the Palomar 60 inch telescope. Due to this, our testing data set is affected by decreased overall throughput (which we currently attempt to compensate for by increased exposure times). It is possible that an increased number of SN Ia spectra can be classified when the mirror has been realuminized. Work on improvements to the SEDM data reduction pipeline is also ongoing (e.g., more effective cosmic-ray rejection and host-galaxy background subtraction).

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Facilities: P48, P60(SEDM).

**Appendix A**

**Loss Function Definitions**

The MSE is the mean of the squared distances between the target variable \( y_i \) and predicted values \( y_i^p \):

\[
\text{MSE} = \frac{1}{n} \sum_{i=1}^{n} (y_i - y_i^p)^2.
\]

(A1)

The MBE is the mean of the distances between \( y_i \) and \( y_i^p \):

\[
\text{MBE} = \frac{1}{n} \sum_{i=1}^{n} (y_i - y_i^p)\cdot
\]

(A2)

The MPE is the mean of the distances between \( y_i \) and \( y_i^p \) divided by \( y_i \):

\[
\text{MPE} = \frac{1}{n} \sum_{i=1}^{n} \frac{(y_i - y_i^p)}{y_i}.
\]

(A3)

The mean absolute error (MAE) is the mean of the absolute distances between \( y_i \) and \( y_i^p \):

\[
\text{MAE} = \frac{1}{n} \sum_{i=1}^{n} |y_i - y_i^p|.
\]

(A4)

**Appendix B**

**SNID and DASH ROC Curves**

To evaluate the performance of SNID (v5.0), we use the standard template bank included with the software\(^20\); the SNe Ia and non-SN templates from the Berkeley SN Ia program (BSNIP; Silverman et al. 2012); the SN Ib and Ic templates from Modjaz et al. (2014, 2016), Liu et al. (2016), and Williamson et al. (2019); and the SN IIP templates from Gutiérrez et al. (2017). The ROC curve shown in Figure 4 was computed by varying the minimal threshold for the rlap parameter between 0 and 25. For each automatic classification we take the best SNID match with the highest rlap value for that particular spectrum, as we have found that this performs better than considering multiple matches (e.g., counting how many SN Ia matches are among the top 10 best matches). The redshift range was restricted to a reasonable range for the SN expected to be found by the BTS (\( z < 0.2 \)). We place no restriction on the phase of the templates, and we have not attempted to restrict the template set used when running SNID. For future work it may be of interest to investigate if there is an optimal set of templates to use for SN Ia binary classification.

In order to evaluate DASH (v1.0 with Models_v06), we used the default template bank included with the software.\(^21\) To create the ROC curve we have investigated the effect of cuts on both the softmax scores and the rlap values that DASH produces. We investigate the ranges softmax > 0.2–1 and rlap > 0–25. To create the ROC curve shown in Figure 4 we consider all possible cuts in the range softmax > 0.5–0.9 and rlap > 0–25, because for this range we observe the most

\(^{20}\) https://people.lam.fr/blondin.stephane/software/snid/

\(^{21}\) https://github.com/daniel-muthukrishna/astodash
stable behavior in DASH. More extreme cuts on $\text{softmax} > 0.92–0.99$ can offer some improvement on the BTS18 data set, but only for one specific and narrow range of FPR (FPR = 0.02), which then gives TPR = 0.62. The rest of the ROC curve becomes noisy and with no real improvement over the one produced for $\text{softmax} > 0.5–0.9$. As such, we consider this behavior to be unstable and do not recommend extreme cuts on the softmax score for SEDM data.

Appendix C
Classification Performance Depending on Data Quality

It is of interest to investigate how $\text{SNIascore}$ performs as a function of data quality. For SN classification, the S/N of a spectrum on its own is not an ideal tracer, because S/N cannot separate between SN-dominated and host-galaxy-dominated spectra. Instead, we use the $r_{\text{lap}}$ values produced by SNID, which correlate positively with S/N and negatively with host-galaxy contamination. To create a high-S/N sample, we use our testing data set (Table 1) and select spectra with $r_{\text{lap}} \geq 0.8$. To create a low-S/N sample, we select spectra with $0 < r_{\text{lap}} < 8$.

The performance of $\text{SNIascore}$ on these high- and low-S/N samples is shown in Figure 6. There is a clear correlation between S/N and $\text{SNIascore}$. For real SN Ia spectra, the scores for low-S/N spectra peak at 0.95. The scores for high-S/N SN Ia spectra peak at 1, with the majority of high-S/N SN Ia spectra receiving scores higher than 0.95. The number of false positives at high $\text{SNIascore}$ does not decrease in the high-S/N sample compared to the low-S/N sample. Selecting only high-S/N spectra is thus unlikely to eliminate the most likely false positives discussed in Section 6.

Most importantly, we find that very low FPR can be achieved regardless of the overall S/N of the sample and that the same cut to minimize FPR ($\text{SNIascore} > 0.8$), found in Section 5.1 remains appropriate. $\text{SNIascore}$ has successfully learned to consistently give low scores to bad-quality data. This finding is important because ground-based observations are always subject to bad observing conditions.

It is unnecessary to apply any external cuts based on S/N before running $\text{SNIascore}$ on the data SEDM is producing that pass the quality checks of the data reduction pipeline (Blagorodnova et al. 2018; Rigault et al. 2019). The main impact of conditions leading to data of bad quality is that fewer spectra out of the total number of spectra that are observed can be successfully classified.

Figure 6. Classification performance of $\text{SNIascore}$ on high-S/N ($r_{\text{lap}} \geq 0.8$) and low-S/N ($0 < r_{\text{lap}} < 8$) subsamples selected from the testing data set. Top-left panel: the $\text{SNIascore}$ distributions for both “Real SNe Ia” spectra (red) and spectra of “Other” transients (blue). The high-S/N data set is shown as solid bars, and the low-S/N data set is shown as open bars. $N_{\text{tot}}$ is the total for each class separately (i.e., the sum of the “Real SNe Ia” bins add up to unity). Bottom panel: FPR and FNR as a function of $\text{SNIascore}$ for the high-S/N (solid red and blue lines) and for the low-S/N (dashed red and blue lines) data sets. A cut at $\text{SNIascore} > 0.8$ results in an FPR of zero on both the high- and low-S/N data sets. Right panel: ROC curve for $\text{SNIascore}$ on the high-S/N data set (red line) and the low-S/N data set (blue line).
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