Automated Seismic Source Characterisation Using Deep
Graph Neural Networks

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Key Points:

• We propose a deep learning approach for automated earthquake location and mag-
nitude estimation based on Graph Neural Network theory
• This new approach processes multi-station waveforms and incorporates station lo-
cations explicitly
• Including station locations improves the accuracy of epicentre estimation compared
to models that are location-agnostic

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Abstract

Most seismological analysis methods require knowledge of the geographic location of the stations comprising a seismic network. However, common machine learning tools used in seismology do not account for this spatial information, and so there is an underutilised potential for improving the performance of machine learning models. In this work, we propose a Graph Neural Network (GNN) approach that explicitly incorporates and leverages spatial information for the task of seismic source characterisation (specifically, location and magnitude estimation), based on multi-station waveform recordings. Even using a modestly-sized GNN, we achieve model prediction accuracy that outperforms methods that are agnostic to station locations. Moreover, the proposed method is flexible to the number of seismic stations included in the analysis, and is invariant to the order in which the stations are arranged, which opens up new applications in the automation of seismological tasks and in earthquake early warning systems.

Plain language summary

To determine the location and size of earthquakes, seismologists use the geographic locations of the seismic stations that record the ground shaking in their data analysis workflow. By taking the distance between stations and the relative timing of the onset of the shaking, the origin of the seismic waves can be accurately reconstructed. In recent years, machine learning (a subfield of artificial intelligence) has shown great potential to automate seismological tasks, such as earthquake source localisation. Most machine learning methods do not take into consideration the geographic locations of the seismic stations, and so the usefulness of these methods could still be improved by providing the locations at which the data was recorded. In this work, we propose a method that accounts for geographic locations of the seismic stations, and we show that this improves the machine learning predictions.

1 Introduction

Seismic source characterisation is a primary task in earthquake seismology, and involves the estimation of the epicentral location, hypocentral depth, and moment of the seismic source. Particularly for the purposes of earthquake early warning, emergency response and timely information dissemination, an estimate of the seismic source characteristics needs to be produced rapidly, preferably without the intervention of an analyst.
One computational tool that satisfies these requirements is machine learning, making it a potential candidate to address the challenge of rapid seismic source characterisation.

Recently, attempts have been made to apply machine learning to seismic source characterisation (Käufl et al., 2014; Perol et al., 2018; Lomax et al., 2019; Kriegerowski et al., 2019; Mousavi & Beroza, 2020b,a). In the ConvNetQuake approach of Perol et al. (2018), a convolutional neural network was adopted to distinguish between noise and earthquake waveforms, and to determine the regional earthquake cluster from which each event originated. This method was extended by Lomax et al. (2019) to global seismicity. Mousavi & Beroza (2020b) employed a combined convolutional-recurrent neural network to estimate earthquake magnitudes. It is noteworthy that these methods only accept single-station waveforms as an input, which goes against the common intuition that at least three seismic stations are required to triangulate and locate a seismic source. One possible explanation for the performance of these methods is that they rely on waveform similarity (Perol et al., 2018) and differences in phase arrival times (Mousavi & Beroza, 2020b). Unfortunately, since the parametrisation through high-dimensional machine learning methods does not carry a clear physical meaning, this hypothesis is not easily tested.

Alternatively, a multi-station approach would take as input for each earthquake all the waveforms recorded by the seismic network. One compelling argument in favour of single-station approaches is that for each earthquake there are as many training samples as there are stations, whereas in the multi-station approach there is only one training sample per earthquake (the concatenated waveforms from the whole network). Since the performance of a deep learning model tends to benefit from larger volumes of data available for training, the model predictions may not improve when combining multiple station data into a single training sample. Second, micro-earthquakes are usually not recorded on multiple seismic stations if the seismic network is sparse, warranting further development of single-station methods. Lastly, concatenating data from multiple stations in a meaningful way is non-trivial. If the seismic network has a Euclidean structure, i.e. if it is arranged in a regular pattern like for uniformly-spaced seismic arrays or fibre-optic distributed acoustic sensing, the data can be naturally arranged into e.g. a 2D image, where the distance between each pixel is representative of the spatial sampling distance. Unfortunately, most seismic networks are not arranged in a regular structure, so that the geometry of the network needs to be learned implicitly, as was attempted by Kriegerowski et al. (2019). Even though this approach yielded acceptable hypocentre location estimates,
it remains an open question whether better results could be achieved when the non-Euclidean nature of the seismic network is better accounted for. Moreover, the seismic stations comprising the network may not be continuously operational over the period of interest (due to (de)commissioning, maintenance, or temporary campaigning strategies), leading to gaps in the fixed Euclidean data structure. Rather, seismic networks are better represented by a time-varying graph structure.

The deep learning tools most commonly used in seismology, convolutional neural networks (CNNs) and multi-layer perceptrons (MLPs) (see also Supplementary Text S1; Rosenblatt, 1957; Fukushima, 1980; Rumelhart et al., 1986; LeCun et al., 2015; Schramowski et al., 2020), are well suited to Euclidean data structures, but are not optimal for graph data structures. One important characteristic of graphs is that they are not defined by the ordering or positioning of the data, but only by the relations between data. As such, valid operations on a graph need to be invariant to the data order. This is not generally the case for CNNs, which exploit ordering as a proxy for spatial distance, nor for MLPs, which rely on the constant structure of the input features. Fortunately, much progress has been made in the field of Graph Neural Networks (GNNs; Gori et al., 2005; Scarselli et al., 2009; Zhou et al., 2019), providing a robust framework for analysing non-Euclidean data using existing deep learning tools.

In this contribution, we will demonstrate how GNNs can be applied to seismic source characterisation using data from multiple seismic stations simultaneously. The method does not require a fixed seismic network configuration, and so the number of stations to be included in each sample is allowed to vary over time. Moreover, the stations do not need to be ordered geographically or as a function of distance from the seismic source. This makes the proposed method suitable for earthquake early warning and disaster response applications, in which the number and location of stations on which a given event is recorded is not known a-priori.

2 Methods

2.1 Basic Concepts of Graph Neural Networks

Over the past several years, numerous deep learning techniques have been proposed that allow for the analysis of non-Euclidean data structures (Bronstein et al., 2017; Zhou et al., 2019), which has found applications in point cloud data (Qi et al., 2017; Wang et
al., 2019), curved manifolds (Monti et al., 2017), and $N$-body classical mechanics (Sanchez-Gonzalez et al., 2019), among many others. As a subclass of non-Euclidean objects, graphs highlight relations between objects, typically represented as nodes connected by edges. Commonly studied examples of graph-representable objects include social networks (Hamilton et al., 2017), molecules (Duvenaud et al., 2015), and urban infrastructures (Cui et al., 2019). Owing to the lack of spatial ordering of graph structures, mathematical operations performed on graphs need to be invariant to the order in which the operations are executed. Moreover, nodes and relations between them (i.e. the edges) may not be fixed, and so the graph operations need to generalise to an arbitrary number of nodes and/or edges (and potentially the number of graphs) at any given moment. In essence, suitable graph operations are those that can be applied to the elements of a set of unknown cardinality. These can be simple mathematical operations such as taking the mean, maximum, or sum of the set, or they can involve more expressive aggregation (Battaglia et al., 2018) and message passing (Gilmer et al., 2017) operations.

To make the above statement more concrete, we represent a seismic network by an edgeless graph in which each seismic station is a node. In the context of seismic source characterisation, information travels from the seismic source to each individual receiver station independently of the relative positions between the stations. Since no information is transmitted from one station to another, it is not intuitive to include e.g. the relative distance between two stations. While local site amplifications could play an important role in the seismic source characterisation process, such information should be encoded in the absolute location of each station rather than the relative location. Hence, for the task of seismic source characterisation, the relations between individual stations are not physically meaningful, and so we do not include edges connecting the nodes in the analysis, reducing the graph to an unordered set. While a graph with no edges may seem ludicrous, the existence of edges is not a requirement for defining a graph, and basic architectural principles (e.g. Battaglia et al., 2018) still apply. Naturally, in cases where the relation between stations is relevant, for example in seismic array beamforming (which relies on relative locations and arrival times), edge information should be included. Each node in our graph carries two attributes: a three-component seismic waveform time-series, and a geographic location. The graph itself carries four attributes: the latitude, longitude, depth, and magnitude of the seismic source. Through suitable processing and ag-
Figure 1. Synoptic overview of the adopted model architecture. The three-component waveforms from a receiver station are fed into a CNN, after which the extracted features are combined with the station’s geographic location and further processed by an MLP. The resulting node feature vector of all the stations are aggregated, and this aggregated feature vector is passed through a second MLP that predicts the seismic source characteristics.

2.2 Model architecture

The model architecture employed in this work consists of three components that operate sequentially – see Fig. 1 and Supplementary Text S2 for details (Tompson et al., 2015; Saxe et al., 2014; Hu et al., 2020). Firstly, we analyse the waveforms of a given station using a CNN. This CNN processes the three-component waveform (comprising $N_t$ time samples) and extracts a set of $N_f$ features. The geographic location (latitude/longitude)
of the seismic station is then appended to produce a feature vector of size $N_f + 2$. This feature vector serves as an input for the second component: an MLP that recombines the time-series features and station location into a final station-specific feature vector of size $N_q$. This process is repeated for all $N_s$ stations in the network using the same CNN and MLP components (i.e. the exact same operations are applied to each station individually). The convolution operations are performed only along the time axis. The output of the CNN after concatenation with each station location is then of size $N_s \times (N_f + 2)$, and the output of the MLP is of size $N_s \times N_q$.

After processing of the node attributes (the waveforms and locations of each station), the output of the MLP is max reduced over all stations to yield a graph feature vector. Empirically we have found that a max reduce yields better results than averaging or summation. The extracted features carry no physical meaning, and the information content of the feature vectors adapts to the type of aggregation during training. Hence, the most suitable type of aggregation needs to be determined experimentally. Finally, the graph feature vector is fed into a second MLP to predict the graph attributes, being the latitude, longitude, depth, and magnitude of the seismic source. Each of these source attributes is scaled so that they fall within the continuous range of $-1 < x < +1$, enforced by a tanh activation function in the last layer in the network. In contrast to previous work (Perol et al., 2018; Lomax et al., 2019), no binning of the source characteristics is performed. Moreover, we do not perform event detection, as this has already been done in numerous previous studies (Dysart & Pulli, 1990; Li et al., 2018; Mousavi et al., 2019; Wu et al., 2019, and others) and is essentially a solved problem. Instead, we focus on the characterisation of a given seismic event. Note that the procedure above is intrinsically invariant to the number and ordering of the seismic stations: the feature extraction and re-combination with the geographic location is performed for each node individually and does not incorporate information from the other stations in the network. The aggregation and the resulting graph feature vector are also independent of the number and ordering of stations. Finally, the seismic source characteristics are predicted from this invariant graph feature vector, and are hence completely independent of the network input ordering and size.

To regularise the learning process, we include dropout regularisation (Srivastava et al., 2014) with a dropout rate of 15\% between each layer in each model component. Since the mechanics of convolutional layers are different from “dense” layers (those defin-
ing the MLPs), we use spatial dropout regularisation (Tompson et al., 2015) that randomly sets entire feature maps of a convolutional layer to zero (as opposed to individual elements in the feature maps). The use of dropout regularisation is dually motivated: first of all it reduces overfitting on the training set, as the model cannot rely on a single layer output (which could be randomly set to zero), promoting redundancy and generalisation within the model. Secondly, by randomly perturbing the data flow within the neural networks, the model output becomes probabilistic. The probability distribution of the model predictions for a given event can be acquired by evaluating a given input multiple times at inference time, with the variability produced by the dropout regularisation. This technique is commonly referred to as Bayesian dropout (Gal & Ghahramani, 2016), as it yields a posterior distribution and hence provides a means to estimate the epistemic uncertainty for the predictions.

2.3 Data description and training procedure

To construct a training set, we use ObsPy (Beyreuther et al., 2010) to download the broadband station inventory and earthquake catalogue of the Southern California Seismic Network (SCSN; Hutton et al., 2010) over the period 2000-2015. For both the seismic station and event locations, we limit the latitude range from 32° to 36°, and the longitude range from −120° to −116°. The lower earthquake magnitude limit is set to 3 with no depth cut-off. In total, 1377 events and 187 stations are included in the data set. After downloading the three-component waveforms and removing the instrument response, we filter the waveforms to a 0.1-8 Hz bandpass and interpolate onto a common time base of 1 ≤ t ≤ 101 seconds after the event origin time, over 2048 evenly spaced time samples (≈ 20 Hz sampling frequency). For an average P-wave speed of 6 km s⁻¹, this time interval allows the stations at the far ends of the domain (roughly 440×440 km in size) to record the event while keeping the data volume compact. The lower limit of the frequency band is chosen below the corner frequency of the earthquakes in this analysis (MW < 6, with corresponding corner frequency fc > 0.2 Hz; Madariaga, 1976) such that information regarding the seismic moment is retained. The upper frequency limit acknowledges the common notion that attenuation and scattering rapidly reduce the signal spectrum at higher frequencies. Although the start time of all selected waveforms is fixed relative to their event origin time, the shift-equivariance of the convolution layers ensures that the extracted features are not sensitive to their timing with re-
spect to the origin. Subsequent aggregation over the time-axis renders the features strictly
time-invariant. As a result, selecting a different start of the data time window (which
is inevitable when the event origin time is unknown) does not affect the model perfor-
mane. The processed waveforms are then scaled by their standard deviation and stored
in a database which includes the locations of the seismic stations that have recorded the
events. Note that not all stations are operational at the time of a given event, and hence
the number of stations with recordings of the event varies.

After processing the waveforms, the locations of the stations and seismic source are
scaled by the minimum and maximum latitude/longitude, so that the re-scaled locations
fall in the range of ±1. Such normalisation is generally considered good practice in deep
learning. Similarly, the source depth is scaled to fall in the same range by taking a min-
imum and maximum source depth of 0 and 30 km respectively. The earthquake magni-
tude is scaled taking a minimum and maximum of 3 and 6. The full data set is then ran-
domly split 80-20 into a training set and a validation set, respectively. A batch of train-
ing samples is generated on the fly between training epochs by randomly selecting 16 train-
ing events, and 50 randomly selected stations associated with each event, which we con-
sider to strike a good balance between data volume and memory consumption. When
a given event was recorded by fewer than 50 stations, the absent recordings are replaced
by zeros (which do not contribute to the model performance). The model performance
is evaluated through a mean absolute error loss between the predicted and target seis-
mic source characteristics (scaled between ±1), and training is performed by minimisa-
tion of the loss using the ADAM algorithm (Kingma & Ba, 2017). Training is contin-
ued for 500 epochs, at which point the model performance has saturated. On a single
nVidia Tesla K80, the training phase took about 1 hour in total. Once trained, evalu-
ation of 1377 events with up to 50 stations each takes less than 5 s of computation time
(including data transfer overhead), or 3.5 ms per event.

3 Results and Discussion

3.1 Reference model performance

We evaluate the performance of the trained model on both the training and val-
idation data sets separately (Fig. 2a-e and Supplementary Figure S2). The model pos-
terior is estimated by maintaining dropout regularisation at inference time (as discussed
**Figure 2.** (a)-(e) Prediction error distributions for the trained model, for (a) latitude, (b) longitude, (c) epicentre, (d) depth, and (e) magnitude of each event. The model performance when including the station geographic locations is evaluated separately for the train and validation data sets, showing minimal overfitting. When the station locations are omitted, the performance is evaluated on the combined data set; (f) Residuals of the epicentral locations. Each arrow represents one catalogued event, starting at the predicted epicentre and pointing towards the catalogue epicentre. The colours indicate the ratio of the misfit over the 95% confidence interval of the model posterior. Hence, blue colours indicate that the catalogue epicentre falls within the 95% confidence interval, and red colours that the epicentre falls outside of it; (g) Overlay of the locations of seismic stations on the interpolated prediction error (in km).
in the previous section), and performing the inference 100 times on each event in the training and validation catalogues and calculating the corresponding mean and standard deviation. Overall, the performance is similar for either data set, which indicates that over-fitting on the training set is minimal. The mean absolute difference between the catalogue values and the model predictions is less than 0.11° (≈ 13 km in distance) for the latitude and longitude (which amounts to a mean epicentral location error of 18 km), 3.3 km for the depth, and 0.13 for the event magnitude. While these predictions are not as precise as typical non-relocated estimates for Southern California (Powers & Jordan, 2010), they are obtained without phase picking or waveform amplitude modelling, nor is a crustal velocity models explicitly provided (though it is implicitly encoded in the catalogue hypocentre locations). Hence, the method provides a reasonable first-order estimate of location and magnitude that can serve as a starting point for subsequent refinement based on traditional seismological tools.

Since we can compute the posterior distribution for each event, we can compare the confidence intervals given by the posterior with the true epicentre location error. In Fig. 2f, we plot the residual vectors between the predicted epicentre locations and those in the catalogue. To visualise the model uncertainty, we compute an error ratio metric as the distance between the predicted and catalogued epicentres, normalised by the 95% confidence interval obtained from the model posterior. Hence, values less than 1 indicate that the true epicentre location falls within the 95% confidence interval, while values greater than 1 indicate the converse. Most of the predictions have an error ratio < 1. This assessment of the uncertainty in the predictions only addresses epistemic uncertainties, but does not immediately address aleatoric uncertainties (errors or bias on the SCSN catalogue). The epicentral errors reported for the SCSN catalogue are approximately 2 km, even though an in-depth analysis of these errors suggests that this error assessment is somewhat over-estimated (Powers & Jordan, 2010). The expected aleatoric uncertainties are therefore much smaller than the epistemic uncertainties given by the model posterior distribution.

The spatially interpolated prediction error seems partly correlated with the local density of seismic stations (Fig. 2g), as regions with the highest station density also exhibit a low prediction error. The largest systematic errors are found in the northwest and southeast corners of the selected domain, where the station density is low and where the model seems unable to achieve the bounding values of latitude and longitude. This ob-
observation can be explained by the behaviour of the tanh activation function, which asymptotically approaches its range of $\pm 1$, corresponding with the range of latitudes and longitudes of the training samples. Hence, increasingly larger activations are required to push the final location predictions towards the boundaries of the domain, biasing the results towards the interior. This highlights a fundamental trade-off between resolution (prediction accuracy) in the interior of the data domain, and the maximum amplitude of the predictions (which also applies to linear activation functions).

Lastly, we perform additional analyses of the sensitivity of the predictions to the signal-to-noise ratio, waveform pre-processing, and epicentre location (Supplementary Figures S4-S6). These analyses show that the predictions are rather robust to the event magnitude (as a proxy for signal-to-noise ratio), and insensitive to instrument corrections. Moreover, preliminary tests, in which we adopted a filter passband of 0.5-5 Hz, indicated that the choice for the pre-filtering frequency band had little influence on the model performance. When the model is provided with waveforms belonging to an event with an epicentre outside of the selected training domain, the model predictions for the epicentre location collapse to an average value around the centre of the domain (Supplementary Figure S6). Fortunately, the uncertainty of the predictions (inferred from the posterior distribution of each event) is also much larger than for events that are located within the domain. Thus, exterior events can be distinguished from interior events through the inferred precision.

### 3.2 Influence of geographic information on location accuracy

A direct test to assess whether the station geographic location information is actually used in making the predictions (and therefore holds predictive value), we perform inference on the full data set, but set the station coordinates to a fixed mean value of $(34^\circ, -118^\circ)$ – see Fig. 2a-e and Supplementary Figure S3. While the predictions for the event magnitude remain mostly unchanged, the estimation of the epicentre location deteriorates and becomes broadly distributed (typical for random predictions). This clearly indicates that the station location information plays an important role in estimating the epicentre locations. Thus, the adopted GNN approach, in which station location information is provided explicitly, holds an advantage over station-location agnostic methods. Interestingly, the event magnitude is almost as well resolved as when the station coordinates are included, which suggests that the model relies on the waveform data but
not on station locations to estimate the magnitude. This was also observed by Mousavi & Beroza (2020b), who proposed that the relative timing of the P- and S-wave arrivals may encode epicentral distance information. Combined with the amplitude of the waveforms, this may implicitly encode magnitude information.

Related to this, we investigate the effect of the (maximum) number of stations included at inference time by selecting, for each event, the stations recording the waveforms with the $M$ highest standard deviations. All other waveforms are set to zero and therefore do not contribute to the predictions. If a given event was recorded by fewer than $M$ stations, only the maximum number of operational stations was used with no augmentation. We perform the inference for $M = \{1, 2, 5, 10, 15, 20, 30, 40, 50\}$ stations, and compute the mean absolute error of the predictions for the epicentre location (expressed
as a distance in km; Fig. 3a), hypocentral depth (Fig. 3b), and event magnitude (Fig. 3c).

For all the predicted quantities, we observe that the misfit with the catalogue values rapidly decreases with the maximum number of stations included in the analysis, until the performance saturates at around $M \geq 40$. The reason for this saturation may lie in the distribution of the number of operational stations per event (Fig. 3d). Since the majority of catalogued events is recorded by fewer than 40 stations, increasing $M$ beyond 40 is only potentially beneficial only for a small number of events. For reference, we compute two performance baselines: firstly, we take the mean value of each quantity (latitude, longitude, depth, magnitude) over the catalogue and calculate the mean absolute error relative to these. This baseline represents the performance of a “biased coin flip” (i.e. random guessing). Secondly, we train our model specifically using only a single station per training sample, through which the method specialises to single-waveform analysis (c.f. Perol et al., 2018; Lomax et al., 2019; Mousavi & Beroza, 2020b). These baselines are included in Fig. 3 as horizontal dotted and dashed-dotted lines for the mean absolute error relative to the (constant value) mean, and for the single-station model, respectively. Strikingly, the model that was trained on the single-station waveforms achieves worse performance in terms of the predicted hypocentre locations than the model trained on 50 stations, but using only a single station at inference time. A possible explanation for this, is that the single-station model may have gotten attracted to a poor local minimum in the loss landscape, after which the model started over-fitting, whereas the 50-station model was able to generalise better and descended into a better local minimum.

Lastly, we compare our model performance with a model that treats the seismic network as an Euclidean object, and hence has no explicit knowledge of the geographic locations of the seismic stations (“station-location agnostic”). This station-location agnostic model only features components #1 and #3 (see Fig. 1 and Supplementary Text S3 for details) and does not incorporate the station locations among the data features. Instead, the stations appear in a fixed order in a matrix of size $N_s \times N_t \times 3$, where $N_s = 256$ denotes the total number of stations in the network (187) plus zero padding to make $N_s$ an integer power of two. Potentially, the station-location agnostic model is able to “learn” the configuration of the seismic network and implicitly utilise station locations in predicting the seismic source characteristics. As in most traditional CNN approaches, we use a 2D kernel of size $k_s \times k_t$ with $k_s = 3$ so that information from “neighbouring” stations (i.e. sequentially appearing in the grid, which does not imply geographic
proximity) is combined into the next layer of the model. Downsampling of the data is performed along both the temporal and station axes. Even though the number of free parameters of the station-location agnostic model is almost twice that of the graph-based model (owing to the larger convolutional kernels), and even though the model has access to all the stations simultaneously, the prediction error of the seismic source parameters is significantly larger (dashed line in Fig. 3). Moreover, the station-location agnostic model required 5 times more computation time per training epoch. Hence, the GNN approach proposed here offers substantial benefits in terms of predictive power and ease of training.

3.3 Potential applications

The method proposed in this study does not require the intervention of an analyst to prepare or verify the model input data (e.g. picking P- and S-wave first arrivals), and so it can operate autonomously. This, combined with the rapid inference time of \( \approx 3.5 \) ms for 50 stations, opens up applications in automated source characterisation that require a rapid response, such as earthquake early warning (EEW; Allen & Melgar, 2019), emergency response, and timely public dissemination. The aim of this study is to demonstrate the potential of incorporating seismic station locations (and possibly other node or edge attributes in a graph structure). Therefore, the model architecture was not optimised with the purpose of EEW in mind. Nonetheless, its modular nature allows for modifications required to accommodate the real-time demands of EEW.

The first out of three components of this model consists of a CNN that analyses the waveforms of each seismic station and yields a set of station-specific features. The advantage of using a CNN is that it has immediate access to all the available information to produce a set of features optimal for the subsequent MLP components. Alternatively, a different class of deep neural networks suitable for time-series analysis, the Recurrent Neural Networks (RNN; Hochreiter & Schmidhuber, 1997; Sherstinsky, 2020), allows for online (real-time) processing of time series. Within the generalised framework of GNNs (Battaglia et al., 2018), replacing the first CNN component with an RNN produces an equally valid model architecture, still independent of the number and ordering of stations. As such, for each new data entry the model updates its prediction, taking into account previously seen data (the “memory” of the RNN). A robust prediction will be one for which the output of the model converges to a stable estimate of hypocen-
tre location and magnitude. Since we here employed a CNN rather than an RNN, we do not know how much time since the first ground motions is required to converge to a stable prediction, and we anticipate that this convergence depends on the quality and consistency of the data. Moreover, different components of the prediction may converge at different rates: while the hypocentre estimate may be governed by the (first) arrival of seismic energy at the various stations in the region (and therefore on the station density), the magnitude estimate is potentially controlled by the duration of the moment-rate function (Meier et al., 2017). Owing to the opacity of our deep learning method, we cannot directly assess which part of the input governs which part in the output, and so this will need to be assessed empirically.

As mentioned in Section 2.2, we focussed our efforts on seismic source characterisation and not event detection. For any EEW task, earthquake detection is a crucial first step, which fortunately has been demonstrated to be a task suitable for machine learning methods (e.g. Dysart & Pulli, 1990; Li et al., 2018; Mousavi et al., 2019; Wu et al., 2019). In the methods proposed in the present study, earthquake detection could be performed by adding an additional graph attribute (alongside latitude, longitude, depth, and magnitude) indicating whether or not an event has been detected (similar to Perol et al., 2018; Lomax et al., 2019). Alternatively, a dedicated detection algorithm (based on machine learning or otherwise), could run in parallel and trigger the source characterisation algorithm once an event has been detected. This second approach significantly reduces computational overhead. Flexibility in the number of stations included in the model input facilitates processing of an expanding data set as more seismic stations experience ground shaking after the first detection.

For the applications of emergency response and information dissemination, the real-time requirements are less stringent, so that some response time may be sacrificed in favour of prediction accuracy, maintaining the CNN component #1. Our method can be readily applied to automated earthquake catalogue generation in regions where large volumes of raw data exist, but which have not been fully processed. This typically arises in aftershock campaigns with stations that were not telemetered, for instance Ocean Bottom Seismometers. Given the relatively small size of the GNN employed here, re-training a pre-trained model on data from a different region is relatively inexpensive. Out of the 110,836 trainable parameters, less than half (42,244) reside in the second and third components of the network. The first CNN component is completely agnostic to any spa-
tional or regional information, as it only extracts features from time series of individual sta-
tions. Hence, if the waveforms in the target region are similar to those in the initial train-
ing region, the first component requires no re-training. This leaves only the smaller sec-
ond and third MLP components to be re-trained and adapted to the characteristics of
the target region. As such, fewer training seismic events than employed for the initial
training will be required for fine-tuning of the model. It is crucial to realise here that
the second and third components potentially encode the crustal velocity structure and
local site amplifications, and are therefore specific to the domain that was selected dur-
ing training (Southern California). Direct application of the trained model to other re-
gions without retraining is unwarranted. The scaling of the re-trained model performance
with the number of stations will need to be assessed empirically, as it may be sensitive
to station redundancy, and spatial coverage and density.

Aside from automatically providing an earthquake catalogue, the estimates of the
seismic source locations can offer a suitable starting point for additional seismological
analyses. With the re-trained model, the predicted hypocentre locations yield approx-
imate phase arrival times at the various stations in the seismic network, which serve as
a basis to set the windows for cross-correlation time-delay estimation and subsequent double-
difference relocation. Grid-search based inversion efforts could be directed to a region
around the predicted hypocentre location, rather than expanding the search of candi-
date source locations to a much larger (regional) domain. Even though the model pre-
dictions for the epicentral locations are larger than what conventional seismological tech-
niques can achieve, there is merit in deep-learning based automated source character-
isation to expedite current seismological workflows.

Lastly, we point out that the GNN-approach is rather general, and that it may be
adopted in other applications such as seismic event detection or classification, that ben-
efit from geographic or relational information of the seismic network. Aside from pre-
predicting “global" graph attributes, like was done in this study, GNNs can also be employed
to predict node or edge attributes. Examples of such attributes include site amplifica-
tion factors and event detections for the nodes (seismic stations), and phase associations
for the edges. Since many geophysical data are inherently non-Euclidean, graph-based
approaches offer a natural choice for the analysis of these data, and permit creative so-
lutions to present-day challenges.
4 Conclusions

In this study we propose a method to incorporate the geometry of a seismic network into deep learning architectures using a Graph Neural Network (GNN) approach, applied to the task of seismic source characterisation (earthquake location and magnitude estimation). By incorporating the geographic location of stations into the learning and prediction process, we find that the deep learning model achieves superior performance in predicting the seismic source characteristics (epicentral latitude/longitude, hypocentral depth, and event magnitude) compared to a model that is agnostic to the layout of the seismic network. In this way, multi-station waveforms can be incorporated while preserving flexibility to the number of available seismic stations, and invariance to the ordering of the station recordings. The GNN-based approach warrants the exploration of new avenues in earthquake early warning and rapid earthquake information dissemination, as well as in automated earthquake catalogue generation or other seismological tasks.

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