

Automatic change-point detection in time series via deep learning

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Abstract

Detecting change points in data is challenging because of the range of possible types of change and types of behaviour of data when there is no change. Statistically efficient methods for detecting a change will depend on both of these features, and it can be difficult for a practitioner to develop an appropriate detection method for their application of interest. We show how to automatically generate new offline detection methods based on training a neural network. Our approach is motivated by many existing tests for the presence of a change point being representable by a simple neural network, and thus a neural network trained with sufficient data should have performance at least as good as these methods. We present theory that quantifies the error rate for such an approach, and how it depends on the amount of training data. Empirical results show that, even with limited training data, its performance is competitive with the standard cumulative sum (CUSUM) based classifier for detecting a change in mean when the noise is independent and Gaussian, and can substantially outperform it in the presence of auto-correlated or heavy-tailed noise. Our method also shows strong results in detecting and localizing changes in activity based on accelerometer data.

Keywords: automatic statistician, classification, likelihood-free inference, neural networks, structural breaks, supervised learning

1 Introduction

Detecting change points in data sequences is of interest in many application areas such as bioinformatics (Picard et al., 2005), climatology (Reeves et al., 2007), signal processing (Haynes et al., 2017), and neuroscience (Oh et al., 2005). In this work, we are primarily concerned with the problem of offline change-point detection, where the entire data is available to the analyst beforehand. Over the past few decades, various methodologies have been extensively studied in this area, see Killick et al. (2012), Jandhyala et al. (2013), Fryzlewicz (2014, 2023), Wang and Samworth (2018), Truong et al. (2020) and references therein. Most research on change-point detection has concentrated on detecting and localizing different types of change, e.g. change in mean (Fryzlewicz, 2014; Killick et al., 2012), variance (Gao et al., 2019; Li et al., 2015), median (Fryzlewicz, 2021), or slope (Baranowski et al., 2019; Fearnhead et al., 2019), amongst many others.

Many change-point detection methods are based upon modelling data when there is no change and when there is a single change, and then constructing an appropriate test statistic to detect the presence of a change (e.g. Fearnhead & Rigaill, 2020; James et al., 1987). The form of a good test statistic will vary with our modelling assumptions and the type of change we wish to detect. This can lead to difficulties in practice. As we use new models, it is unlikely

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that there will be a change-point detection method specifically designed for our modelling assumptions. Furthermore, developing an appropriate method under a complex model may be challenging, while in some applications an appropriate model for the data may be unclear but we may have substantial historical data that shows what patterns of data to expect when there is, or is not, a change.

In these scenarios, currently a practitioner would need to choose the existing change detection method which seems the most appropriate for the type of data they have and the type of change they wish to detect. To obtain reliable performance, they would then need to adapt its implementation, for example tuning the choice of threshold for detecting a change. Often, this would involve applying the method to simulated or historical data.

To address the challenge of automatically developing new change detection methods, this paper is motivated by the question: Can we construct new test statistics for detecting a change based only on having labelled examples of change points? We show that this is indeed possible by training a neural network to classify whether or not a dataset has a change of interest. This turns changepoint detection in a supervised learning problem.

A key motivation for our approach are results that show many common test statistics for detecting changes, such as the CUSUM test for detecting a change in mean, can be represented by simple neural networks. This means that with sufficient training data, the classifier learnt by such a neural network will give performance at least as good as classifiers corresponding to these standard tests. In scenarios where a standard test, such as CUSUM, is being applied but its modelling assumptions do not hold, we can expect the classifier learnt by the neural network to outperform it.

There has been increasing recent interest in whether ideas from machine learning, and methods for classification, can be used for change-point detection. Within computer science and engineering, these include a number of methods designed for and that show promise on specific applications (e.g. Ahmadzadeh, 2018; De Ryck et al., 2021; Gupta et al., 2022; Huang et al., 2023). Within statistics, Londschien et al. (2022) and Lee et al. (2023) consider training a classifier as a way to estimate the likelihood-ratio statistic for a change. However, these methods train the classifier in an unsupervised way on the data being analysed, using the idea that a classifier would more easily distinguish between two segments of data if they are separated by a change point. Chang et al. (2019) use simulated data to help tune a kernel-based change-detection method. Methods that use historical, labelled data have been used to train the tuning parameters of change-point algorithms (e.g. Hocking et al., 2015; Liehrmann et al., 2021). Also, neural networks have been employed to construct similarity scores of new observations to learned pre-change distributions for online changepoint detection (Lee et al., 2023). However, we are unaware of any previous work using historical, labelled data to develop offline change-point methods. As such, and for simplicity, we focus on the most fundamental aspect, namely the problem of detecting a single change. Detecting and localizing multiple changes is considered in Section 6 when analysing activity data. We remark that by viewing the change-point detection problem as a classification instead of a testing problem, we aim to control the overall mis-classification error rate (MER) instead of handling the Type I and Type II errors separately. In practice, asymmetric treatment of the two error types can be achieved by suitably reweighting mis-classification in the two directions in the training loss function.

The method we develop has parallels with likelihood-free inference methods (Beaumont, 2019; Gourieroux et al., 1993) in that one application of our work is to use the ability to simulate from a model so as to circumvent the need to analytically calculate likelihoods. However, the approach we take is very different from standard likelihood-free methods which tend to use simulation to estimate the likelihood function itself. By comparison, we directly target learning a function of the data that can discriminate between instances that do or do not contain a change (though see Gutmann et al., 2018 for likelihood-free methods based on re-casting the likelihood as a classification problem).

For an introduction to the statistical aspects of neural network-based classification, albeit not specifically in a change-point context, see Ripley (1994).

We now briefly introduce our notation. For any $n \in \mathbb{Z}^+$, we define $[n] := \{1, ..., n\}$. We take all vectors to be column vectors unless otherwise stated. Let $\mathbf{1}_n$ be the all-one vector of length n. Let $\mathbb{I}_{\{\cdot\}}$ represent the indicator function. The vertical symbol $|\cdot|$ represents the absolute value or cardinality of \cdot depending on the context. For vector $\mathbf{x} = (x_1, ..., x_n)^{\mathsf{T}}$, we define its p-norm as $\|\mathbf{x}\|_p := (\sum_{i=1}^n |x_i|^p)^{1/p}, p \ge 1$; when $p = \infty$, define $\|\mathbf{x}\|_{\infty} := \max_i |x_i|$. All proofs, as well as additional simulations and real data analyses appear in the online supplementary material.



Figure 1. A neural network with two hidden layers and width vector m = (4, 4).

2 Neural networks

The initial focus of our work is on the binary classification problem for whether a change point exists in a given time series. We will work with multi-layer neural networks with Rectified Linear Unit (ReLU) activation functions and binary output. The multi-layer neural network consists of an input layer, hidden layers, and an output layer, and can be represented by a directed acyclic graph, see Figure 1. Let $L \in \mathbb{Z}^+$ represent the number of hidden layers and $m = (m_1, \ldots, m_L)^{\mathsf{T}}$ the vector of the hidden layer widths, i.e. m_i is the number of nodes in the *i*th hidden layer. For a neural network with L hidden layers, we use the convention that $m_0 = n$ and $m_{L+1} = 1$. For any bias vector $\boldsymbol{b} = (b_1, b_2, \ldots, b_r)^{\mathsf{T}} \in \mathbb{R}^r$, define the shifted activation function $\sigma_{\boldsymbol{b}}: \mathbb{R}^r \to \mathbb{R}^r$:

$$\sigma_{\boldsymbol{b}}((y_1, \ldots, y_r)^{\mathsf{T}}) = (\sigma(y_1 - b_1), \ldots, \sigma(y_r - b_r))^{\mathsf{T}},$$

where $\sigma(x) = \max(x, 0)$ is the ReLU activation function. The neural network can be mathematically represented by the composite function $h : \mathbb{R}^n \to \{0, 1\}$ as

$$h(\mathbf{x}) := \sigma_{\lambda}^* W_L \sigma_{b_L} W_{L-1} \sigma_{b_{l-1}} \cdots W_1 \sigma_{b_1} W_0 \mathbf{x}, \tag{1}$$

where $\sigma_{\lambda}^{*}(x) = \mathbb{1}\{x > \lambda\}, \lambda > 0$ and $W_{\ell} \in \mathbb{R}^{m_{\ell+1} \times m_{\ell}}$ for $\ell \in \{0, \dots, L\}$ represent the weight matrices. We define the function class $\mathcal{H}_{L,m}$ to be the class of functions h(x) with *L* hidden layers and width vector *m*.

The output layer in equation (1) employs the shifted heaviside function $\sigma_i^*(x)$, which is used for binary classification as the final activation function. This choice is guided by the fact that we use the 0-1 loss, which focuses on the percentage of samples assigned to the correct class, a natural performance criterion for binary classification. Besides its wide adoption in machine learning practice, another advantage of using the 0–1 loss is that it is possible to utilise the theory of the Vapnik– Chervonenkis (VC) dimension (see, e.g. Shalev-Shwartz & Ben-David, 2014, Definition 6.5) to bound the generalization error of a binary classifier equipped with this loss; indeed, this is the approach we take in this work. The relevant results regarding the VC dimension of neural network classifiers are, e.g. in Bartlett et al. (2019). As in Schmidt-Hieber (2020), we work with the exact minimizer of the empirical risk. In both binary or multi-class classification, it is possible to work with other losses which make it computationally easier to minimise the corresponding risk, see, e.g. Bos and Schmidt-Hieber (2022), who use a version of the cross-entropy loss. However, loss functions different from the 0-1 loss make it impossible to use VC-dimension arguments to control the generalization error, and more involved arguments, such as those using the covering number (Bos & Schmidt-Hieber, 2022) need to be used instead. We do not pursue these generalizations in the current work.

3 CUSUM-based classifier and its generalizations are neural networks

3.1 Change in mean

We initially consider the case of a single change point with an unknown location $\tau \in [n-1], n \ge 2$, in the model

$$\begin{split} &X=\boldsymbol{\mu}+\boldsymbol{\xi},\\ &\boldsymbol{\mu}=(\mu_{\mathrm{L}}\mathbb{1}\{i\leq\tau\}+\mu_{\mathrm{R}}\mathbb{1}\{i>\tau\})_{i\in[n]}\in\mathbb{R}^{n}, \end{split}$$

where $\mu_{\rm L}$, $\mu_{\rm R}$ are the unknown signal values before and after the change point; $\boldsymbol{\xi} \sim N_n(0, I_n)$. The CUSUM test is widely used to detect mean changes in univariate data. For the observation \boldsymbol{x} , the CUSUM transformation $\mathcal{C}: \mathbb{R}^n \to \mathbb{R}^{n-1}$ is defined as $\mathcal{C}(\boldsymbol{x}) := (\boldsymbol{v}_1^{\mathsf{T}}\boldsymbol{x}, \ldots, \boldsymbol{v}_{n-1}^{\mathsf{T}}\boldsymbol{x})^{\mathsf{T}}$, where $\boldsymbol{v}_i := (\sqrt{\frac{n-i}{in}}\mathbf{1}_{n-i}^{\mathsf{T}})^{\mathsf{T}}$ for $i \in [n-1]$. Here, for each $i \in [n-1], (\boldsymbol{v}_i^{\mathsf{T}}\boldsymbol{x})^2$ is the log likelihood-ratio statistic for testing a change at time *i* against the null of no change (e.g. Baranowski et al., 2019). For a given threshold $\lambda > 0$, the classical CUSUM test for a change in the mean of the data is defined as

$$b_{\lambda}^{\text{CUSUM}}(\boldsymbol{x}) = \mathbb{1}\{\|\mathcal{C}(\boldsymbol{x})\|_{\infty} > \lambda\}.$$

The following lemma shows that $h_i^{\text{CUSUM}}(x)$ can be represented as a neural network.

Lemma 3.1 For any $\lambda > 0$, we have $h_{\lambda}^{\text{CUSUM}}(x) \in \mathcal{H}_{1,2n-2}$.

The fact that the widely used CUSUM statistic can be viewed as a simple neural network has far-reaching consequences: this means that given enough training data, a neural network architecture that permits the CUSUM-based classifier as its special case cannot do worse than CUSUM in classifying change-point vs. no-change-point signals. This serves as the main motivation for our work, and a prelude to our next results.

3.2 Beyond the mean change model

We can generalise the simple change in mean model to allow for different types of change or for non-independent noise. In this section, we consider change-point models that can be expressed as a change in regression problem, where the model for data given a change at τ is of the form

$$X = Z\beta + c_\tau \phi + \Gamma \xi, \tag{2}$$

where for some $p \ge 1$, Z is an $n \times p$ matrix of covariates for the model with no change, c_{τ} is an $n \times 1$ vector of covariates specific to the change at τ , and the parameters β and ϕ are, respectively, a $p \times 1$ vector and a scalar. The noise is defined in terms of an $n \times n$ matrix Γ and an $n \times 1$ vector of independent standard normal random variables, ξ .

For example, the change in mean problem has p = 1, with Z a column vector of ones, and c_{τ} being a vector whose first τ entries are zeros, and the remaining entries are ones. In this formulation, β is the pre-change mean and ϕ is the size of the change. The change in slope problem (Fearnhead et al., 2019) has p = 2 with the columns of Z being a vector of ones, and a vector whose *i*th entry is *i*; and c_{τ} has *i*th entry that is max{0, $i - \tau$ }. In this formulation, β defines the pre-change linear mean and ϕ the size of the change in slope. Choosing Γ to be proportional to the identity matrix gives a model with independent, identically distributed noise; but other choices would allow for auto-correlation.

The following result is a generalization of Lemma 3.1, which shows that the likelihood-ratio test for equation (2), viewed as a classifier, can be represented by our neural network.

Lemma 3.2 Consider the change-point model (2) with a possible change at $\tau \in [n-1]$. Assume further that Γ is invertible. Then there is an $h^* \in \mathcal{H}_{1,2n-2}$ equivalent to the likelihood-ratio test for testing $\phi = 0$ against $\phi \neq 0$. Importantly, this result shows that for this much wider class of change-point models, we can replicate the likelihood-ratio-based classifier for change using a simple neural network.

Other types of changes can be handled by suitably pre-transforming the data. For instance, squaring the input data would be helpful in detecting changes in the variance and if the data followed an AR(1) structure, then changes in auto-correlation could be handled by including transformations of the original input of the form $(x_t x_{t+1})_{t=1,...,n-1}$. On the other hand, even if such transformations are not supplied as the input, a neural network of suitable depth is able to approximate these transformations and consequently successfully detect the change (Schmidt-Hieber, 2020, Lemma A.2). This is illustrated in Figure S3 of the online supplementary material, where we compare the performance of neural network-based classifiers of various depths constructed with and without using the transformed data as inputs.

4 Generalization error of neural network change-point classifiers

In Section 3, we showed that CUSUM and generalised CUSUM could be represented by a neural network. Therefore, with a large enough amount of training data, a trained neural network classifier that included CUSUM, or generalised CUSUM, as a special case, would perform no worse than it on unseen data. In this section, we provide generalization bounds for a neural network classifier for the change-in-mean problem, given a finite amount of training data. En route to this main result, stated in Theorem 4.3, we provide generalization bounds for the CUSUM-based classifier, in which the threshold has been chosen on a finite training dataset.

We write $P(n, \tau, \mu_L, \mu_R)$ for the distribution of the multivariate normal random vector $X \sim N_n(\mu, I_n)$, where $\mu := (\mu_L \mathbb{1}\{i \le \tau\} + \mu_R \mathbb{1}\{i > \tau\})_{i \in [n]}$. Define $\eta := \tau/n$. Lemma 4.1 and Corollary 4.1 control the mis-classification error of the CUSUM-based classifier.

Lemma 4.1 Fix $\varepsilon \in (0, 1)$. Suppose $X \sim P(n, \tau, \mu_L, \mu_R)$ for some $\tau \in \mathbb{Z}^+$ and $\mu_L, \mu_R \in \mathbb{R}$.

(a) If
$$\mu_{L} = \mu_{R}$$
, then $\mathbb{P}\{\|\mathcal{C}(X)\|_{\infty} > \sqrt{2\log(n/\varepsilon)}\} \le \varepsilon$.
(b) If $\|\mu_{L} - \mu_{R}\|\sqrt{\eta(1-\eta)} > \sqrt{8\log(n/\varepsilon)/n}$, then $\mathbb{P}\{\|\mathcal{C}(X)\|_{\infty} \le \sqrt{2\log(n/\varepsilon)}\}$
 $\le \varepsilon$.

For any B > 0, define

$$\Theta(B) := \left\{ (\tau, \mu_{\rm L}, \mu_{\rm R}) \in [n-1] \times \mathbb{R} \times \mathbb{R} : |\mu_{\rm L} - \mu_{\rm R}| \sqrt{\tau(n-\tau)}/n \in \{0\} \cup (B, \infty) \right\}.$$

Here, $|\mu_L - \mu_R| \sqrt{\tau(n-\tau)}/n = |\mu_L - \mu_R| \sqrt{\eta(1-\eta)}$ can be interpreted as the signal-to-noise ratio (SNR) of the mean change problem. Thus, $\Theta(B)$ is the parameter space of data distributions where there is either no change or a single change point in mean whose SNR is at least *B*. The following corollary controls the mis-classification risk of a CUSUM statistics-based classifier:

Corollary 4.1 Fix B > 0. Let π_0 be any prior distribution on $\Theta(B)$, then draw $(\tau, \mu_L, \mu_R) \sim \pi_0$ and $X \sim P(n, \tau, \mu_L, \mu_R)$, and define $Y = \mathbb{I}\{\mu_L \neq \mu_R\}$. For $\lambda = B\sqrt{n/2}$, the classifier $b_{\lambda}^{\text{CUSUM}}$ satisfies

$$\mathbb{P}(h_1^{\text{CUSUM}}(X) \neq Y) \le ne^{-nB^2/8}.$$

Theorem 4.2 below, which is based on Corollary 4.1, Bartlett et al. (2019, Theorem 7) and Mohri et al. (2012, Corollary 3.4), shows that the empirical risk minimizer in the neural network class $\mathcal{H}_{1,2n-2}$ has good generalization properties over the class of change-point problems parameterised by $\Theta(B)$. Given training data ($X^{(1)}, Y^{(1)}$), ..., ($X^{(N)}, Y^{(N)}$) and any $h : \mathbb{R}^n \to \{0, 1\}$, we define the empirical risk of h as

$$L_N(h) := \frac{1}{N} \sum_{i=1}^N \mathbb{1}\{Y^{(i)} \neq h(\mathbf{X}^{(i)})\}.$$



Figure 2. Plot of the test set mis-classification error rate, computed on a test set of size $N_{\text{test}} = 30,000$, against training sample size N for detecting the existence of a change point on data series of length n = 100. We compare the performance of the CUSUM test and neural networks from four function classes: $\mathcal{H}_{1,m^{(1)}}$, $\mathcal{H}_{1,m^{(2)}}$, $\mathcal{H}_{5,m^{(1)}\mathbf{1}_5}$, and $\mathcal{H}_{10,m^{(1)}\mathbf{1}_0}$, where $m^{(1)} = 4\lfloor \log_2(n) \rfloor$ and $m^{(2)} = 2n - 2$, respectively, under scenarios S1, S1 ', S2, and S3 described in Section 5. (a) Scenario S1 with $\rho_t = 0$. (b) Scenario S1 ' with $\rho_t = 0.7$. (c) Scenario S2 with $\rho_t \sim \text{Unif}([0, 1])$. (d) Scenario S3 with Cauchy noise.

Theorem 4.2 Fix B > 0 and let π_0 be any prior distribution on $\Theta(B)$. We draw $(\tau, \mu_L, \mu_R) \sim \pi_0, X \sim P(n, \tau, \mu_L, \mu_R)$, and set $Y = \mathbb{1}\{\mu_L \neq \mu_R\}$. Suppose that the training data $\mathcal{D} := ((X^{(1)}, Y^{(1)}), \ldots, (X^{(N)}, Y^{(N)}))$ consist of independent copies of (X, Y) and $b_{\text{ERM}} := \arg \min_{h \in \mathcal{H}_{1,2n-2}} L_N(h)$ is the empirical risk minimizer. There exists a universal constant C > 0 such that for any $\delta \in (0, 1)$, equation (3) holds with probability $1 - \delta$.

$$\mathbb{P}(h_{\text{ERM}}(\mathbf{X}) \neq Y \mid \mathcal{D}) \le ne^{-nB^2/8} + C\sqrt{\frac{n^2 \log(n) \log(N) + \log(1/\delta)}{N}}.$$
 (3)

The theoretical results derived for the neural network-based classifier, here and below, all rely on the fact that the training and test data are drawn from the same distribution. However, we observe that in practice, even when the training and test sets have different error distributions, neural network-based classifiers still provide accurate results on the test set; see our discussion of Figure 2 in Section 5 for more details. The mis-classification error in equation (3) is bounded by two terms. The first term represents the mis-classification error of CUSUM-based classifier, see Corollary 4.1, and the second term depends on the complexity of the neural network class measured in its VC dimension. Theorem 4.2 suggests that for training sample size $N \gg n^2 \log n$, a well-trained single hidden layer neural network with 2n - 2 hidden nodes would have comparable performance to that of the CUSUM-based classifier. However, as we will see in Section 5, in practice, a much smaller training sample size N is needed for the neural network to be competitive in the changepoint detection task. This is because the 2n - 2 hidden layer nodes in the neural network representation of $h_{\lambda}^{\text{CUSUM}}$ encode the components of the CUSUM transformation $(\pm v_t^{\top}x:t \in [n-1])$, which are highly correlated.

By suitably pruning the hidden layer nodes, we can show that a single hidden layer neural network with $O(\log n)$ hidden nodes is able to represent a modified version of the CUSUM-based classifier with essentially the same mis-classification error. More precisely, let $Q := \lfloor \log_2 (n/2) \rfloor$ and write $T_0 := \{2^q : 0 \le q \le Q\} \cup \{n - 2^q : 0 \le q \le Q\}$. We can then define

$$b_{\lambda^*}^{\text{CUSUM}_*}(X) = \mathbb{1}l\{\max_{t\in T_0} | \boldsymbol{v}_t^{\mathsf{T}} X| > \lambda^*\}.$$

By the same argument as in Lemma 3.1, we can show that $h_{\lambda^*}^{\text{CUSUM}_*} \in \mathcal{H}_{1,4\lfloor \log_2(n) \rfloor}$ for any $\lambda^* > 0$. The following theorem shows that high classification accuracy can be achieved under a weaker training sample size condition compared to Theorem 4.2.

Theorem 4.3 Fix B > 0 and let the training data \mathcal{D} be generated as in Theorem 4.2. Let $h_{\text{ERM}} := \arg \min_{h \in \mathcal{H}_{L,m}} L_N(h)$ be the empirical risk minimizer for a neural network with $L \ge 1$ layers and $\boldsymbol{m} = (m_1, \ldots, m_L)^{\top}$ hidden layer widths. If $m_1 \ge 4 \lfloor \log_2(n) \rfloor$ and $m_r m_{r+1} = O(n \log n)$ for all $r \in [L-1]$, then there exists a universal constant C > 0 such that for any $\delta \in (0, 1)$, equation (4) holds with probability $1 - \delta$.

$$\mathbb{P}(h_{\text{ERM}}(X) \neq Y \mid \mathcal{D}) \le 2\lfloor \log_2(n) \rfloor e^{-nB^2/24} + C \sqrt{\frac{L^2 n \log^2(Ln) \log(N) + \log(1/\delta)}{N}}.$$
(4)

Theorem 4.3 generalises the single hidden layer neural network representation in Theorem 4.2 to multiple hidden layers. In practice, multiple hidden layers help to keep the MER low even when N is small, see Section 5. Theorems 4.2 and 4.3 are examples of how to derive generalization errors of a neural network-based classifier in the change-point detection task. The same workflow can be employed in other types of changes, provided that suitable representation results of likelihood-based tests in terms of neural networks (e.g. Lemma 3.2) can be obtained. In a general result of this type, the generalization error of the neural network will again be bounded by a sum of the error of the likelihood-based classifier together with a term originating from the VC-dimension bound of the complexity of the neural network architecture.

We further remark that for simplicity of discussion, we have focused our attention on data models where the noise vector $\xi = X - \mathbb{E}X$ has independent and identically distributed normal components. However, since CUSUM-based tests are available for temporally correlated or sub-Weibull data, with suitably adjusted test threshold values, the above theoretical results readily generalise to such settings. See Theorems S4 and S6 in the online supplementary material for more details.

5 Numerical study

We now investigate empirically our approach of learning a change-point detection method by training a neural network. Motivated by the results from the previous section, we will fit a neural network with a single layer and consider how varying the number of hidden layers and the amount of training data affects performance. We will compare to a test based on the CUSUM statistic, both for scenarios where the noise is independent and Gaussian, and for scenarios where there is auto-correlation or heavy-tailed noise. The CUSUM test can be sensitive to the choice of threshold, particularly when we do not have independent Gaussian noise, so we tune its threshold based on training data.

When training the neural network, we first standardise the data onto [0, 1], i.e. $\tilde{x}_i = ((x_{ij} - x_i^{\min})/(x_i^{\max} - x_i^{\min}))_{j \in [n]}$, where $x_i^{\max} := \max_j x_{ij}$, $x_i^{\min} := \min_j x_{ij}$. This makes the neural network procedure invariant to either adding a constant to the data or scaling the data by a constant, which are natural properties to require. We train the neural network by minimizing the cross-entropy loss on the training data. We run training for 200 epochs with a batch size of 32 and a learning rate of 0.001 using the Adam optimizer (Kingma & Ba, 2015). These hyperparameters are chosen based on a training dataset with cross-validation, more details can be found in Section 2 of the online supplementary material.

We generate our data as follows. Given a sequence of length *n*, we draw $\tau \sim \text{Unif}\{2, \dots, n-2\}$, set $\mu_{\text{L}} = 0$ and draw $\mu_{\text{R}} | \tau \sim \text{Unif}([-1.5b, -0.5b] \cup [0.5b, 1.5b])$, where $b := \sqrt{\frac{8n \log(20n)}{\tau(n-\tau)}}$ is chosen

in line with Lemma 4.1 to ensure a good range of SNRs. We then generate $x_1 = (\mu_L \mathbb{1}_{\{t \le \tau\}} + \mu_R \mathbb{1}_{\{t > \tau\}} + \varepsilon_t)_{t \in [n]}$, with the noise $(\varepsilon_t)_{t \in [n]}$ following an AR(1) model with possibly time-varying auto-correlation $\varepsilon_t | \rho_t = \zeta_1$ for t = 1 and $\rho_t \varepsilon_{t-1} + \zeta_t$ for $t \ge 2$, where $(\zeta_t)_{t \in [n]}$ are independent, possibly heavy-tailed noise. The auto-correlations ρ_t and innovations ζ_t are from one of the four scenarios:

S1: n = 100, $N \in \{100, 200, \dots, 700\}$, $\rho_t = 0$, and $\xi_t \sim N(0, 1)$. S1': n = 100, $N \in \{100, 200, \dots, 700\}$, $\rho_t = 0.7$, and $\xi_t \sim N(0, 1)$. S2: n = 100, $N \in \{100, 200, \dots, 1, 000\}$, $\rho_t \sim \text{Unif}([0, 1])$, and $\xi_t \sim N(0, 2)$. S3: n = 100, $N \in \{100, 200, \dots, 1, 000\}$, $\rho_t = 0$, and $\xi_t \sim \text{Cauchy}(0, 0.3)$.

The above procedure is then repeated N/2 times to generate independent sequences $x_1, \ldots, x_{N/2}$ with a single change, and the associated labels are $(y_1, \ldots, y_{N/2})^T = \mathbf{1}_{N/2}$. We then repeat the process another N/2 times with $\mu_{\rm R} = \mu_{\rm L}$ to generate sequences without changes $x_{N/2+1}, \ldots, x_N$ with $(y_{N/2+1}, \ldots, y_N)^{\mathsf{T}} = \mathbf{0}_{N/2}$. The data with and without change $(x_i, y_i)_{i \in [N]}$ are combined and randomly shuffled to form the training data. The test data are generated in a similar way, with a sample size $N_{\text{test}} = 30,000$ and the slight modification that $\mu_{\text{R}} | \tau \sim$ Unif $([-1.75b, -0.25b] \cup [0.25b, 1.75b])$ when a change occurs. We note that the test data is drawn from the same distribution as the training set, though potentially having changes with SNRs outside the range covered by the training set. We have also conducted robustness studies to investigate the effect of training the neural networks on scenario S1 and test on S1', S2, or S3. Qualitatively similar results to Figure 2 have been obtained in this mis-specified setting (see Figure S2 of the online supplementary material). We compare the performance of the CUSUM-based classifier with the threshold cross-validated on the training data with neural networks from four function classes: $\mathcal{H}_{1,m^{(1)}}$, $\mathcal{H}_{1,m^{(2)}}$, $\mathcal{H}_{5,m^{(1)}1_5}$ and $\mathcal{H}_{10,m^{(1)}1_{10}}$, where $m^{(1)} =$ $4\lfloor \log_2(n) \rfloor$ and $m^{(2)} = 2n - 2$, respectively (cf. Theorem 4.3 and Lemma 3.1). Figure 2 shows the test MER of the four procedures in the four scenarios S1, S1', S2, and S3. We observe that when data are generated with independent Gaussian noise (Figure 2a), the trained neural networks with $m^{(1)}$ and $m^{(2)}$ single hidden layer nodes attain very similar test MER compared to the CUSUM-based classifier. This is in line with our Theorem 4.3. More interestingly, when noise has either auto-correlation (Figure 2b and c) or heavy-tailed distribution (Figure 2d), trained neural networks with (L, \mathbf{m}) : $(1, m^{(1)})$, $(1, m^{(2)})$, $(5, m^{(1)}\mathbf{1}_5)$, and $(10, m^{(1)}\mathbf{1}_{10})$ outperform the CUSUM-based classifier, even after we have optimised the threshold choice of the latter. In addition, as shown in Figure S1 in the online supplementary material, when the first two layers of the network are set to carry out truncation, which can be seen as a composition of two ReLU operations, the resulting neural network outperforms the Wilcoxon statistics-based classifier (Dehling et al., 2015), which is a standard benchmark for change-point detection in the presence of heavytailed noise. Furthermore, from Figure 2, we see that increasing L can significantly reduce the average MER when $N \leq 200$. Theoretically, as the number of layers L increases, the neural network is better able to approximate the optimal decision boundary, but it becomes increasingly difficult to train the weights due to issues such as vanishing gradients (He et al., 2016). A combination of these considerations leads us to develop deep neural network architecture with residual connections for detecting multiple changes and multiple change types in Section 6.

6 Detecting multiple changes and multiple change types—case study

From the previous section, we see that single and multiple hidden layer neural networks can represent CUSUM or generalised CUSUM tests and may perform better than likelihood-based test statistics when the model is mis-specified. This prompted us to seek a general network architecture that can detect, and even classify, multiple types of change. Motivated by the similarities between signal processing and image recognition, we employed a deep convolutional neural network (CNN) (Yamashita et al., 2018) to learn the various features of multiple change types. However, stacking more CNN layers cannot guarantee a better network because of vanishing gradients in training (He et al., 2016). Therefore, we adopted the residual block structure (He et al., 2016) for our neural network architecture. After experimenting with various architectures with

	Weak SNR			Strong SNR		
	LR ^{oracle}	LR ^{adapt}	NN	LR ^{oracle}	LR ^{adapt}	NN
Class 1	0.9787	0.9457	0.8062	0.9787	0.9341	0.9651
Class 2	0.8443	0.8164	0.8882	1.0000	0.7784	0.9860
Class 3	0.8350	0.8291	0.8585	0.9902	0.9902	0.9705
Class 4	0.9960	0.9453	0.8826	0.9980	0.9372	0.9312
Class 5	0.8729	0.8604	0.8353	0.9958	0.9917	0.9147
Accuracy	0.9056	0.8796	0.8660	0.9924	0.9260	0.9672

Table 1. Test classification accuracy of oracle likelihood-ratio-based method (LR^{oracle}), adaptive likelihood-ratio method (LR^{adapt}), and our residual neural network (NN) classifier for set-ups with weak and strong signal-to-noise ratios (SNRs)

Note. Data are generated as a mixture of no change point in mean or variance (Class 1), change in mean only (Class 2), change in variance only (Class 3), no-change in a non-zero slope (Class 4), and change in slope only (Class 5). We report the true positive rate of each class and the accuracy in the last row.

different numbers of residual blocks and fully connected layers on synthetic data, we arrived at a network architecture with 21 residual blocks followed by a number of fully connected layers. Figure S5 of the online supplementary material shows an overview of the architecture of the final general-purpose deep neural network for change-point detection. The precise architecture and training methodology of this network \widehat{NN} can be found in Section 3 of the online supplementary material. Neural architecture search approaches (see Paaß & Giesselbach, 2023, Section 2.4.3) offer principled ways of selecting neural architectures. Some of these approaches could be made applicable in our setting.

We demonstrate the power of our general purpose change-point detection network in a numerical study. We train the network on N = 10,000 instances of data sequences generated from a mixture of no change point in mean or variance, change in mean only, change in variance only, no-change in a non-zero slope, and change in slope only, and compare its classification performance on a test set of size 2,500 against that of oracle likelihood-based classifiers (where we prespecify whether we are testing for change in mean, variance or slope) and adaptive likelihood-based classifiers (where we combine likelihood-based tests using the Bayesian information criterion). Details of the data-generating mechanism and classifiers can be found in Section 2 of the online supplementary material. The classification accuracy of the three approaches in weak and strong SNR settings is reported in Table 1. We see that the neural network-based approach achieves similar classification accuracy as adaptive likelihood-based method for strong SNR. We would not expect the neural network to outperform the oracle likelihood-based classifiers, as it has no knowledge of the exact change type of each time series.

We now consider an application to detecting different types of change. The HASC (Human Activity Sensing Consortium) project data contain motion sensor measurements during a sequence of human activities, including 'stay', 'walk', 'jog', 'skip', 'stair up', and 'stair down'. Complex changes in sensor signals occur during transition from one activity to the next (see Figure 3). We have 28 labels in HASC data, see Figure S6 of the online supplementary material. To agree with the dimension of the output, we drop two dense layers 'Dense(10)' and 'Dense(20)' in Figure S5 of the online supplementary material. The resulting network can be effectively applied for change-point detection in sensory signals of human activities and can achieve high accuracy in change-point classification tasks (Figure S8 of the online supplementary material).

Finally, we remark that our neural network-based change-point detector can be utilised to detect multiple change points. Algorithm 1 outlines a general scheme for turning a change-point classifier into a location estimator, where we employ an idea similar to that of MOSUM (Eichinger & Kirch, 2018) and repeatedly apply a classifier ψ to data from a sliding window of size *n*. Here, we require ψ applied to each data segment $X_{[i,i+n)}^*$ to output both the class label $L_i = 0$ or 1 if no change or a change is predicted and the corresponding probability p_i of having a change. In our particular example, for each data segment $X_{[i,i+n)}^*$ of length n = 700, we define $\psi(X_{[i,i+n)}^*) = 0$ if $\widehat{NN}(X_{[i,i+n)}^*)$



Figure 3. The sequence of accelerometer data in x, y, and z axes. From left to right, there are four activities: 'stair down', 'stay', 'stair up', and 'walk', their change points are 990, 1,691, 2,733, respectively marked by black solid lines. The grey rectangles represent the group of 'no-change' with labels: 'stair down', 'stair up', and 'walk'. The red rectangles represent the group of 'one change' with labels: 'stair down \rightarrow stay', 'stay \rightarrow stair up', and 'stair up \rightarrow walk'.

Algorithm 1 Algorithm for change-point localization

Input: new data $x_1^*, \ldots, x_{n^*}^* \in \mathbb{R}^d$, a trained classifier $\psi : \mathbb{R}^{d \times n} \to \{0, 1\}, \gamma > 0$. 1 Form $X_{[i,i+n)}^* := (x_i^*, \ldots, x_{i+n-1})$ and compute $L_i \leftarrow \psi(X_{[i,i+n)}^*)$ for all $i = 1, \ldots, n^* - n + 1$; 2 Compute $\bar{L}_i \leftarrow n^{-1} \sum_{j=i-n+1}^{i} L_j$ for $i = n, \ldots, n^* - n + 1$; 3 Let $\{[s_1, e_1], \ldots, [s_{\hat{\nu}}, e_{\hat{\nu}}]\}$ be the set of all maximal segments such that $\bar{L}_i \ge \gamma$ for all $i \in [s_r, e_r], r \in [\hat{\nu}]$; 4 Compute $\hat{t}_r \leftarrow \arg\max_{i \in [s_r, e_r]} \bar{L}_i$ for all $r \in [\hat{\nu}]$; Output: Estimated change-points $\hat{\tau}_1, \ldots, \hat{\tau}_{\hat{\nu}}$

predicts a class label in {0, 4, 8, 12, 16, 22} (see Figure S6 in the online supplementary material) and 1 otherwise. The thresholding parameter $\gamma \in \mathbb{Z}^+$ is chosen to be 1/2. Figure 4 illustrates the result of multiple change-point detection in HASC data which provides evidence that the trained neural network can detect both the multiple change types and multiple change points.

7 Discussion

Reliable testing for change points and estimating their locations, especially in the presence of multiple change points, other heterogeneities or untidy data, is typically a difficult problem for the applied statistician: they need to understand what type of change is sought, be able to characterise it mathematically, find a satisfactory stochastic model for the data, formulate the appropriate statistic, and fine-tune its parameters. This makes for a long workflow, with scope for errors at its every stage.

In this paper, we showed how a carefully constructed statistical learning framework could automatically take over some of those tasks and perform many of them 'in one go' when provided with examples of labelled data. This turned the change-point detection problem into a supervised learning problem, and meant that the task of learning the appropriate test statistic and fine-tuning its parameters was left to the 'machine' rather than the human user.

The crucial question was that of choosing an appropriate statistical learning framework. The key factor behind our choice of neural networks was the discovery that the traditionally used likelihood-ratio-based change-point detection statistics could be viewed as simple neural networks, which (together with bounds on generalization errors beyond the training set) enabled us to formulate and prove the corresponding learning theory. However, there are a plethora of other excellent predictive frameworks, such as XGBoost, LightGBM or Random Forests (Breiman, 2001; Chen & Guestrin, 2016; Ke et al., 2017) and it would be of interest to establish whether and why they could or could not provide a viable alternative to neural nets here. Furthermore, if we view the neural network as emulating the likelihood-ratio test statistic, in that it will create test statistics for each possible location of a change and then amalgamate these into a single classifier, then we know that test statistics for nearby changes will often be similar. This suggests that imposing some smoothness on the weights of the neural network may be beneficial.



Figure 4. Change-point detection in HASC data. The red vertical lines represent the underlying change points, the blue vertical lines represent the estimated change points. More details on multiple change-point detection can be found in Section 3 of the online supplementary material.

A further challenge is to develop methods that can adapt easily to input data of different sizes, without having to train a different neural network for each input size. For changes in the structure of the mean of the data, it may be possible to use ideas from functional data analysis so that we pre-process the data, with some form of smoothing or imputation, to produce input data of the correct length.

If historical labelled examples of change points, perhaps provided by subject-matter experts (who are not necessarily statisticians) are not available, one question of interest is whether simulation can be used to obtain such labelled examples artificially, based on (say) a single dataset of interest. Such simulated examples would need to come in two flavours: one batch 'likely containing no change points' and the other containing some artificially induced ones. How to simulate reliably in this way is an important problem, which this paper does not solve. Indeed, we can envisage situations in which simulating in this way may be easier than solving the original unsupervised change-point problem involving the single dataset at hand, with the bulk of the difficulty left to the 'machine' at the learning stage when provided with the simulated data.

For situations where there is no historical data, but there are statistical models, one can obtain training data by simulation from the model. In this case, training a neural network to detect a change has similarities with likelihood-free inference methods in that it replaces analytic calculations associated with a model by the ability to simulate from the model. It is of interest whether ideas from that area of statistics can be used here.

The main focus of our work was on testing for a single offline change point, and we treated location estimation and extensions to multiple-change scenarios only superficially, via the heuristics of testing-based estimation in Section 6. Similar extensions can be made to the online setting once the neural network is trained, by retaining the final *n* observations in an online stream in memory and applying our change-point classifier sequentially. One question of interest is whether and how these heuristics can be made more rigorous: equipped with an offline classifier only, how can we translate the theoretical guarantee of this offline classifier to that of the corresponding location estimator or online detection procedure? In addition to this approach, how else can a neural network, however complex, be trained to estimate locations or detect change points sequentially? In our view, these questions merit further work.

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Data availability

The data underlying this article are available at http://hasc.jp/hc2011/index-en.html. The computer code and algorithm are available in Python Package: AutoCPD.

Supplementary material

Supplementary material is available online at Journal of the Royal Statistical Society: Series B.

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