Brain-like replay for continual learning with artificial neural networks

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Abstract

Artificial neural networks suffer from catastrophic forgetting. Unlike humans, when these networks are trained on something new, they rapidly forget what was learned before. In the brain, a mechanism thought to be important for protecting memories is the replay of neuronal activity patterns representing those memories. In artificial neural networks, such memory replay has been implemented in the form of ‘generative replay’, which can successfully prevent catastrophic forgetting in a range of toy examples. Scaling up generative replay to problems with more complex inputs, however, turns out to be challenging. We propose a new, more brain-like variant of replay in which internal or hidden representations are replayed that are generated by the network’s own, context-modulated feedback connections. In contrast to established continual learning methods, our method achieves acceptable performance on the challenging problem of class-incremental learning on natural images without relying on stored data.

1 Introduction

Artificial neural networks (ANNs) are very bad at retaining old information. When trained on something new, these networks typically rapidly and almost fully forget previously acquired skills or knowledge, a phenomenon referred to as ‘catastrophic forgetting’ (McCloskey & Cohen, 1989; Ratcliff, 1990; French, 1999). In stark contrast, humans are able to continually accumulate information throughout their lifetime. A brain mechanism thought to underlie this ability is the replay of neuronal activity patterns that represent previous experiences (Wilson & McNaughton, 1994; O’Neill et al., 2010), a process orchestrated by the hippocampus but also observed in the cortex. This raises the question whether adding replay to ANNs could help to protect them from catastrophic forgetting.

A straightforward way to add replay to an ANN is to store data from previous tasks and interleave it with the current task’s training data (Figure 1A; Rebuffi et al., 2017; Lopez-Paz & Ranzato, 2017; Nguyen et al., 2018; Rolnick et al., 2019). Storing data is however undesirable for a number of reasons. Firstly, from a machine learning perspective, it is a disadvantage to have to store data as it is not always possible to do so in practice (e.g., due to safety or privacy concerns) and it is problematic when scaling up to true lifelong learning. Secondly, from a cognitive science perspective, if we hope to use replay in ANNs as a model for replay in the brain (McClelland et al., 1995), relying on stored data is unwanted as it is questionable how the brain could directly store data (e.g., all pixels of an image), while empirically it is clear that human memory is not perfect (Quiroga et al., 2008).

An alternative to storing data is to generate the data to be replayed (Figure 1B; Robins, 1995; Shin et al., 2017). In particular, ‘generative replay’, in which a separate generative model is incrementally trained on the observed data, has been shown to achieve state-of-the-art performance on a range of toy examples (Shin et al., 2017; van de Ven & Tolias, 2018). Moreover, in the most difficult continual learning scenario, when classes must be learned incrementally, generative replay is currently the only method capable of performing well without storing data (van de Ven & Tolias, 2019).

An important potential drawback of generative replay, however, is that scaling it up to more challenging problems has been reported to be problematic (Lesort et al., 2019; Aljundi et al., 2019). As a result, class incremental learning with more complex inputs (e.g., natural images) remains an
open problem in machine learning: acceptable performance on such problems has so far only been achieved by methods that explicitly store data, such as iCaRL (Rebuffi et al., 2017). Here, our goal is to find a scalable and biologically plausible way (i.e., without storing data) to make replay work on such more realistic problems.

2 HOW DOES GENERATIVE REPLAY SCALE TO MORE COMPLEX PROBLEMS?

To test how generative replay scales to problems with more complex inputs, we used the CIFAR-100 dataset (Krizhevsky et al., 2009) split up into 10 tasks (or episodes) with 10 classes each (Figure 2A). It is important to note that this problem can be setup in multiple ways [or according to different “scenarios”]. One option is that the network only needs to learn to solve each individual task, meaning that at test time it is clear from which task an image to be classified is (i.e., the choice is always just between 10 possible classes). This is the task-incremental learning (Task-IL) scenario (van de Ven & Tolias, 2019) or the “multi-head” setup (Farquhar & Gal, 2018). But another, arguably more realistic option is that the network eventually must learn to distinguish between all 100 classes. Set up this way, it becomes a class-incremental learning (Class-IL) problem: the network must learn a 100-way classifier, but only observes 10 classes at a time (van de Ven & Tolias, 2019). This more challenging scenario is also referred to as the “single-head” setup (Farquhar & Gal, 2018).

On these two scenarios of split CIFAR-100, we compared the performance of generative replay (GR; Shin et al., 2017), using a convolutional variational autoencoder (VAE; Kingma & Welling, 2013) as generator, with that of established methods such as elastic weight consolidation (EWC; Kirkpatrick et al., 2017), synaptic intelligence (SI; Zenke et al., 2017) and learning without forgetting (LwF; Li & Hoiem, 2017). As baselines, we also included the naive approach of simply fine-tuning the neural network on each new task (None; can be seen as lower bound) and a network that was always trained using the data of all tasks so far (Joint; can be seen as upper bound). For a fair comparison, all methods used similar-sized networks and the same training protocol (see Appendix for details).

When split CIFAR-100 was performed according to the Task-IL scenario, the methods EWC, SI and LwF prevented catastrophic forgetting almost fully (Figure 2B). The standard version of generative replay, however, failed on this task protocol with natural images even in the easiest scenario. When performed according to the more realistic Class-IL scenario, the split CIFAR-100 protocol became substantially more challenging and all compared methods (i.e., generative replay, EWC, SI and LwF) performed very poorly suffering from severe catastrophic forgetting (Figure 2C).

3 BRAIN-INSPIRED REPLAY

The above results indicate that straight-forward implementations of generative replay break down for more challenging problems. A likely reason is that the quality of the generated inputs that are replayed is just too low (Figure 2D). One possible solution would be to use the recent progress in generative modelling with deep neural networks (Goodfellow et al., 2014; van den Oord et al., 2016; Rezende & Mohamed, 2015) to try to improve the quality of the generator. Although this approach might work to certain extent, an issue is that incrementally training high-quality generative models is very challenging as well (Lesort et al., 2019). Moreover, such a solution would not be very efficient, since high-quality generative models can be computationally very costly to train and to sample from.
Instead, we take the brain as example:

- **Replay-through-feedback:** For current versions of generative replay it has been suggested that the generator, as source of the replay, is reminiscent of the hippocampus and that the main model corresponds to the cortex (Figure 1B; Shin et al., 2017). Although this analogy has some merit, one issue is that it ignores that the hippocampus sits atop of the cortex in the brain’s processing hierarchy (Felleman & Van Essen, 1991). Instead, we propose to merge the generator into the main model, by equipping it with generative backward or feedback connections. The first few layers of the resulting model can then be interpreted as corresponding to the early layers of the visual cortex and the top layers as corresponding to the hippocampus (Figure 3A). We implemented this “replay-through-feedback” model as a VAE with added softmax classification layer to the top layer of its encoder (see Appendix).

- **Conditional replay:** With a standard VAE, it is not possible to intentionally generate examples of a particular class. To enable our network to control what classes to generate, we replaced the standard normal prior over the VAE’s latent variables by a Gaussian mixture with a separate mode for each class (Figure 3B; see Appendix). This makes it possible to generate specific classes by restricting the sampling of the latent variables to their corresponding modes. Additionally, for our replay-through-feedback model, such a multi-modal prior encourages a better separation of the internal representations of different classes, as they no longer all have to be mapped onto a single continuous distribution.

- **Gating based on internal context:** The brain processes stimuli differently depending on the context or the task that must be performed (Kuchibhotla et al., 2017; Watanabe & Sakagami, 2007). Moreover, contextual cues (e.g., odours, sounds) can bias what memories are replayed (Rasch et al., 2007; Rudoy et al., 2009; Bendor & Wilson, 2012). A simple but effective way to achieve context-dependent processing in an ANN is to fully gate (or “inhibit”) a different, randomly selected subset of neurons in each hidden layer depending on which task should be performed. This is the approach of context-dependent gating (Masse et al., 2018). But an important disadvantage is that this technique can only be used when information about context (e.g., task identity) is always available—i.e., also at test time, which is not the case for class-incremental learning. However, we realized that in the Class-IL scenario it is still possible to use context gates in the decoder part of our network by conditioning on the “internal context” (Figure 3C; see Appendix). Because when producing replay, our model controls itself from what class to generate samples, and based on that internal decision the correct subset of neurons can be inhibited during the generative
Figure 3: Our proposed brain-inspired modifications to the standard generative replay framework. (A) Replay-through-feedback. The generator is merged into the main model by equipping it with generative feedback connections (B) Conditional replay. To enable the model to generate specific classes, the standard normal prior is replaced by a Gaussian mixture with a separate mode for each class. (C) Gating based on internal context. For every class to be learned, a different subset of neurons in each layer is inhibited during the generative backward pass. (D) Internal replay. Instead of representations at the input level (e.g., pixel level), hidden or internal representations are replayed.

- Internal replay: Our final modification was to replay representations of previously learned tasks or classes not all the way to the input level (e.g., pixel level), but to replay them “internally” or at the “hidden level” (Figure 3D; see Appendix). The brain is also not thought to replay memories all the way down to the input level. Mental images are for example not propagated to the retina (Pearson et al., 2015). From a machine learning point of view the hope is that generating such internal representations will be substantially easier, since the purpose of the early layers of a neural network is to disentangle the complex input level representations. A likely requirement for this internal replay strategy to work is that there are no or very limited changes to the first few layers that are not being replayed. From a neuroscience perspective this seems realistic, as for example the representations extracted by the brain’s early visual areas are indeed not thought to drastically change in adulthood (Smirnakis et al., 2005; Karmarkar & Dan, 2006). To simulate development, we pre-trained the convolutional layers of our model on CIFAR-10, a dataset containing similar but non-overlapping images compared to CIFAR-100 (Krizhevsky et al., 2009). During the incremental training on CIFAR-100, we then froze those convolutional layers and we replayed only through the fully-connected layers. For a fair comparison, all other methods also used pre-trained convolutional layers.

4 BRAIN-INSPIRED MODIFICATIONS ENABLE GENERATIVE REPLAY TO SCALE

To test the effectiveness of these neuroscience-inspired modifications, we applied the resulting “brain-inspired replay” method on the same scenarios as before while using similar-sized networks. We found that our modifications substantially improved the performance of generative replay. In the Task-IL scenario, brain-inspired replay (BI-R) almost fully mitigated catastrophic forgetting and outperformed EWC, SI and LwF (Figure 2B). In the Class-IL scenario, brain-inspired replay also significantly outperformed the other methods, although its performance still remained substantially under the “upper bound” of always training on the data of all classes so far (Figure 2C). Nevertheless, we are not aware of any continual learning method that performs better on this challenging problem without storing data. Finally, we found that combining our brain-inspired replay approach with SI (BI-R + SI) substantially improved performance further, thereby further closing the gap towards the upper bound of joint training (Figure 2C).

5 DISCUSSION

We proposed a new, brain-inspired variant of generative replay in which internal or hidden representations are replayed that are generated by the network’s own, context-modulated feedback connections. As a machine learning contribution, our method is the first to perform well on the challenging problem of class-incremental learning with natural images without relying on stored data. As a cognitive science contribution, our method provides evidence that replay could indeed be a feasible way for the brain to combat catastrophic forgetting.
AUTHOR CONTRIBUTIONS


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APPENDIX

CODE AVAILABILITY

Detailed, well-documented code that can be used to reproduce or build upon the reported experiments will be available online at https://github.com/GMvandeVen/brain-inspired-replay.

TASK PROTOCOL

The CIFAR-100 dataset (Krizhevsky et al., 2009) was split into ten tasks or episodes, such that each task/episode consisted of ten classes. The original 32x32 pixel RGB-color images were normalized (i.e., each pixel-value was subtracted by the relevant channel-wise mean and divided by the channel-wise standard deviation, with means and standard deviations calculated over all training images), but no other pre-processing or augmentation was performed. The standard training/test-split was used resulting in 50,000 training images (500 per class) and 10,000 test images (100 per class).

NETWORK ARCHITECTURE

For a fair comparison, the same base neural network architecture was used for all compared methods. This base architecture had 5 pre-trained convolutional layers followed by 2 fully-connected layers each containing 2000 nodes with ReLU non-linearities and a softmax output layer.

In the Task-IL scenario the softmax output layer was ‘multi-headed’, meaning that always only the output units of classes in the task under consideration – i.e., either the current task or the replayed task – were active; while in the Class-IL scenario the output layer was ‘single-headed’, meaning that always all output units of the classes encountered so far were active.

The convolutional layers contained 16, 32, 64, 128 and 254 channels. Each layer used a 3x3 kernel, a padding of 1, and a stride of 1 in the first layer (i.e., no downsampling) and a stride of 2 in the other layers (i.e., image-size was halved in each of those layers). All convolutional layers used batch-norm (Ioffe & Szegedy, 2015) followed by a ReLU non-linearity. For the 32x32 RGB pixel images used in this study, these convolutional layers returned 256x2x2=1024 image features. No pooling was used. To simulate development, the convolutional layers were pre-trained on CIFAR-10, which is a dataset containing similar but non-overlapping images and image-classes compared to CIFAR-100 (Krizhevsky et al., 2009). Pretraining was done by training the base neural network to classify the 10 classes of CIFAR-10 for 100 epochs, using the ADAM-optimizer (\(\beta_1 = 0.9, \beta_2 = 0.999\)) with learning rate of 0.0001 and mini-batch size of 256.

TRAINING

The neural networks were sequentially trained on all tasks or episodes of the task protocol, with only access to the data of the current task / episode. Each task was trained for 5000 iterations of stochastic gradient descent using the ADAM-optimizer (\(\beta_1 = 0.9, \beta_2 = 0.999\); Kingma & Ba, 2014) with learning rate of 0.0001 and a mini-batch size of 256.

GENERATIVE REPLAY

For standard generative replay, two models were sequentially trained on all tasks: [1] the main model, for actually solving the tasks, and [2] a separate generative model, for generating inputs representative of previously learned tasks. See Figure 4 for a schematic.

Main model  The main model was a classifier with the base neural network architecture. The loss function used to train this model had two terms: one for the data of the current task and one for the replayed data, with both terms weighted according to how many tasks/episodes the model had seen so far:

\[
L_{\text{total}} = \frac{1}{N_{\text{tasks so far}}} L_{\text{current}} + (1 - \frac{1}{N_{\text{tasks so far}}}) L_{\text{replay}}
\]  

(1)
Figure 4: (A) Incremental training protocol with generative replay. On the first task (or episode), the main model \([M]\) and a separate generator \([G]\) are trained normally. When moving on to a new task, these two trained models first produce the samples to be replayed (see panel B). Those generated samples are then replayed along with the training data of the current task, and both models are trained further on this extended dataset. (B) To produce the samples to be replayed, inputs representative of the previous tasks are first sampled from the trained generator. The generated inputs are then labelled based on the predictions made for them by the trained main model, whereby the labels are not simply the most likely predicted classes but vectors with the predicted probabilities for all possible classes (i.e., distillation is used for the replay).

The loss of the current task was the standard classification loss. Because the replayed samples were labelled with soft targets instead of hard targets, standard classification loss could not be used for the replayed data and distillation loss was used instead (see below).

**Generating a replay sample** The data to be replayed was produced by first sampling inputs from the generative model, after which those generated inputs were presented to the main model and labelled with a target vector containing the class probabilities predicted by that model (Figure 4B). There are however a few subtleties regarding the generation of these replay samples.

Firstly, the samples replayed during task \(T\) were generated by the versions of the generator and main model directly after finishing training on task \(T - 1\). Implementationally, this could be achieved either by temporarily storing a copy of both models after finishing training on each task, or by already generating all samples to be replayed during training on an upcoming task before training on that task is started.

Secondly, as is common for distillation, the target probabilities predicted by the main model with which the generated inputs were labelled were made ‘softer’ by raising the temperature \(T\) of the softmax layer. That is, for an input \(x\) to be replayed during training of task \(K\), the soft targets were given by the vector \(\tilde{y}\) with \(c^{th}\) element equal to:

\[
\tilde{y}_c = p_{\hat{\theta}^{(K-1)}}^T (Y = c | x)
\]  

(2)

where \(\hat{\theta}^{(K-1)}\) is the vector with parameter values after finishing training on task \(K - 1\) and \(p_{\hat{\theta}}^T\) is the conditional probability distribution defined by the neural network with parameters \(\theta\) and with the temperature of the softmax layer raised to \(T\). A typical value for this temperature is 2, which was the value used here.

Thirdly, there were subtle differences between the two continual learning scenarios regarding which output units were active when generating the soft targets for the inputs to be replayed. With the Task-IL scenario, only the output units corresponding to the classes of the task intended to be replayed were active (with the task intended to be replayed randomly selected among the previous tasks), while with the Class-IL scenario the output units of the classes from up to the previously learned task were active (i.e., classes in the current task were inactive and were thus always assigned zero probability).

**Distillation loss** The training objective for replayed data with soft targets was to match the probabilities predicted by the model being trained to these soft targets by minimizing the cross entropy between them. For an input \(x\) labeled with a soft target vector \(\tilde{y}\), the per-sample distillation loss is given by:

\[
\mathcal{L}^D (x, \tilde{y}; \theta) = -T^2 \sum_{c=1}^{N_{\text{classes}}} \tilde{y}_c \log p_{\theta}^T (Y = c | x)
\]  

(3)

with temperature \(T\) again set to 2. The scaling by \(T^2\) ensured that the relative contribution of this objective matched that of a comparable objective with hard targets (Hinton et al., 2015).
Generative model A symmetric variational autoencoder (VAE: Kingma & Welling [2013] Rezende et al. [2014]) was used as generator. The VAE consisted of an encoder network $q_\phi$ mapping an input-vector $x$ to a vector of stochastic latent variables $z$, and a decoder network $p_\psi$ mapping those latent variables $z$ to a reconstructed or decoded input-vector $\hat{x}$. We kept the architecture of both networks similar to the base neural network: the encoder consisted of 5 pre-trained convolutional layers (as described above) followed by 2 fully-connected layers containing 2000 ReLU units and the decoder had 2 fully-connected layers with 2000 ReLU units followed by 5 deconvolutional or transposed convolutional layers (Zeiler et al., 2011). Mirroring the convolutional layers, the deconvolutional layers contained 128, 64, 32, 16 and 3 channels. The first four deconvolutional layers used a 4x4 kernel, a padding of 1 and a stride of 2 (i.e., image-size was doubled in each of those layers), while the final layer used a 3x3 kernel, a padding of 1 and a stride of 1 (i.e., no upsampling). The first four deconvolutional layers used batch-norm followed by a ReLU non-linearity, while the final layer had no non-linearity. The stochastic latent variable layer $z$ had 100 Gaussian units parameterized by mean $\mu(x)$ and standard deviation $\sigma(x)$, the outputs of the encoder network $q_\phi$ given input $x$ and the prior over them was the standard normal distribution.

Typically, the parameters of a VAE (collected here in $\phi$ and $\psi$) are trained by maximizing a variational lower bound on the evidence (or ELBO), which is equivalent to minimizing the following per-sample loss function for an input $x$:

$$L^G(x; \phi, \psi) = E_{z \sim q_\phi(.|x)}[-\log p_\psi(x|z)] + D_{KL}(q_\phi(.|x)||p(.))$$

(4)

whereby $q_\phi(.|x) = \mathcal{N} \left( \mu^{(x)}, \sigma^{(x)} \right)$ is the posterior distribution over the latent variables $z$ defined by the encoder given input $x$, $p(.) = \mathcal{N}(0, I)$ is the prior distribution over the latent variables and $D_{KL}$ is the Kullback-Leibler divergence. With this combination of prior and posterior, it has been shown that the “latent variable regularization term” can be calculated, without having to do estimation, as (Kingma & Welling [2013]):

$$L^{\text{latent}}(x; \phi) = \frac{1}{2} \sum_{j=1}^{100} \left( 1 + \log(\sigma_j^{(x)} \sigma_j^{(x)}) - \mu_j^{(x)} \mu_j^{(x)} - \sigma_j^{(x)} \sigma_j^{(x)} \right)$$

(5)

whereby $\mu_j^{(x)}$ and $\sigma_j^{(x)}$ are the $j$th elements of respectively $\mu(x)$ and $\sigma(x)$. To simplify the “reconstruction term”, we made the output of the decoder network $p_\psi$ deterministic, which is a modification that is common in the VAE literature (Doersch [2016] Gregor et al. [2015]). We redefined the reconstruction term as the expected binary cross entropy between the original and the decoded pixel values:

$$L^{\text{recon}}(x; \phi, \psi) = E_{\epsilon \sim \mathcal{N}(0, I)} \left[ \sum_{p=1}^{N_{\text{pixels}}} x_p \log (\tilde{x}_p) + (1 - x_p) \log (1 - \tilde{x}_p) \right]$$

(6)

whereby $x_p$ was the value of the $p$th pixel of the original input image $x$ and $\tilde{x}_p$ was the value of the $p$th pixel of the decoded image $\tilde{x} = p_\psi \left( \mathcal{N}(z(x)) \right)$ with $z(x) = \mu(x) + \sigma(x) \odot \epsilon$ and $\epsilon \sim \mathcal{N}(0, I)$. Replacing $z(x)$ by $\mu(x) + \sigma(x) \odot \epsilon$ is known as the “reparameterization trick” (Kingma & Welling, 2013) and makes that a Monte Carlo estimate of the expectation in Eq. 6 is differentiable with respect to $\phi$. As is common in the literature, for this estimate we used just a single sample of $\epsilon$ for each datapoint.

To train the generator, the same hyperparameters (i.e., learning rate, optimizer, iterations, batch size) were used as for training the main model. Similar to the main model, the generator was also trained with replay (i.e., $L^{\text{g}}_{\text{total}} = \frac{1}{N_{\text{tasks so far}}} \frac{L^{\text{G}}_{\text{current}}}{\text{current}} + \frac{1}{N_{\text{tasks so far}}} \frac{L^{\text{G}}_{\text{replay}}}{\text{replay}}$).

Brain-inspired replay

With the above described standard generative replay approach as starting point, we made the following four modifications:

Replay-through-Feedback The “replay-through-feedback” (RtF) model was implemented as a symmetric VAE with an added softmax classification layer to the final hidden layer of the encoder.
(i.e., the layer before the latent variable layer; Fig. 3A). Now only one model had to be trained. To train this model, the loss function for the data of the current task had two terms that were simply added: $L_{\text{current}} = L^c + L^G$, whereby $L^c$ was the standard classification loss and $L^G$ was the generative loss (see Eq. 4). For the replayed data, as for our implementation of standard generative replay, the classification term was replaced by the distillation term from Eq. 1. 

$L_{\text{total}}^{\text{RtF}} = \frac{1}{N_{\text{tasks so far}}}L_{\text{current}}^{\text{RtF}} + (1 - \frac{1}{N_{\text{tasks so far}}})L_{\text{replay}}^{\text{RtF}}$.

### Conditional Replay

To enable the network to generate examples of specific classes, we replaced the standard normal prior over the stochastic latent variables $z$ by a Gaussian mixture with a separate mode for each class (Figure 3B):

$$p_X(\cdot) = \sum_{c=1}^{N_{\text{classes}}} p(Y = c)p_X(\cdot|c)$$

with $p_X(\cdot|c) = \mathcal{N}(\mu^c, \sigma^c I)$ for $c = 1, \ldots, N_{\text{classes}}$, whereby $\mu^c$ and $\sigma^c$ are the trainable mean and standard deviation of the mode corresponding to class $c$. $\chi$ is the collection of trainable means and standard deviations of all classes and $p(Y = c) = \text{Categorical} \left( \frac{1}{N_{\text{classes}}} \right)$ is the class-prior. Because of the change in prior distribution, the expression for analytically calculating the latent variable regularization term $L_{\text{latent}}$ in Eq. 5 is no longer valid. But for an input $x$ labeled with a hard target $y$ (i.e., for current task data), because the prior distribution over the latent variables $z$ reduces to the mode corresponding to class $y$, it can be shown that $L_{\text{latent}}$ still has a closed-form expression:

$$L_{\text{latent}}(x, y; \phi, \chi) = D_{KL}(q_\phi (\cdot | x) || p_X(\cdot | y))$$

$$= -\int q_\phi(z|x) \log \frac{q_\phi(z|x)}{p_X(z|y)} dz$$

$$= E_{z \sim q_\phi(\cdot|x)} [-\log q_\phi(z|x)] + E_{z \sim q_\phi(\cdot|x)} [\log p_X(z|y)]$$

$$= \frac{1}{2} \sum_{j=1}^{J} \left( 1 + \log(\sigma^2_j) - \log(\sigma^2_y) - \frac{(\mu_j^x - \mu_j^y)^2}{\sigma^2_j} \right)$$

whereby $J$ is the dimensionality of the latent variables $z$ (i.e., $J = 100$ for our experiments) and $\mu_j^y$ and $\sigma_j^y$ are the $j$th elements of respectively $\mu^y$ and $\sigma^y$. The last equality is based on the following two simplifications:

$$E_{z \sim q_\phi(\cdot|x)} [-\log q_\phi(z|x)] = E_{z \sim \mathcal{N}(\mu^x, \sigma^x I)} [-\log \mathcal{N}(z|\mu^x, \sigma^x I)]$$

$$= E_{z \sim \mathcal{N}(\mu^x, \sigma^x I)} \left[ \frac{1}{2} \sum_{j=1}^{J} \frac{(z_j - \mu_j^x)^2}{\sigma_j^x} \right] + \frac{1}{2} \log \left( 2\pi \sigma_j^x \right)$$

$$= E_{z \sim \mathcal{N}(0, I)} \left[ \frac{1}{2} \sum_{j=1}^{J} \left( \mu_j^x + \sigma_j^x \epsilon_j - \mu_j^x \right)^2 \right] + \frac{1}{2} \log(2\pi) + \frac{1}{2} \sum_{j=1}^{J} \log \left( \sigma_j^x \right)$$

$$= \frac{1}{2} \sum_{j=1}^{J} E_{z \sim \mathcal{N}(0,1)} [\epsilon^2] + \frac{1}{2} \log(2\pi) + \frac{1}{2} \sum_{j=1}^{J} \log \left( \sigma_j^x \right)$$

$$= \frac{1}{2} \sum_{j=1}^{J} \left( 1 + \log(2\pi) + \log \left( \sigma_j^x \right) \right)$$

(9)
\[
E_{z \sim q_{\phi}(.|x)} [\log p_{\chi}(z|y)] = E_{z \sim N(\mu^{(x)}, \sigma^{(x)} I)} \left[ \log N \left( z | \mu^{y}, \sigma^{y2} I \right) \right] \\
= E_{z \sim N(\mu^{(x)}, \sigma^{(x)} I)} \left[ -\frac{1}{2} \sum_{j=1}^{J} \frac{(z_j - \mu_j^{y})^2}{\sigma_j^{y2}} - \frac{1}{2} \log \left( 2\pi \right) \prod_{j=1}^{J} \sigma_j^{y2} \right] - \frac{1}{2} \log \left( (2\pi)^{J} \prod_{j=1}^{J} \sigma_j \right) \\
= E_{e \sim N(0, I)} \left[ -\frac{1}{2} \sum_{j=1}^{J} \frac{(\mu_j^{(x)} - \mu_j^{y})^2}{\sigma_j^{y2}} - \frac{1}{2} \sum_{j=1}^{J} \frac{\sigma_j(x)^2}{\sigma_j^{y2}} + J \log(2\pi) + \sum_{j=1}^{J} \log \left( \sigma_j^{y2} \right) \right] \\
= -\frac{1}{2} \sum_{j=1}^{J} \left( \frac{\mu_j^{(x)} - \mu_j^{y}}{\sigma_j^{y2}} \right)^2 + \frac{\sigma_j(x)^2}{\sigma_j^{y2}} + \log(2\pi) + \sum_{j=1}^{J} \log \left( \sigma_j^{y2} \right) \\
\tag{10}
\]

However, for an input \( \mathbf{x} \) labeled with a soft target \( \hat{y} \) (i.e., for replayed data), because the prior distribution over the latent variables \( z \) is no a Gaussian mixture, there is no closed-form expression for \( L^{\text{latent}} \). We therefore resort to estimation by sampling, for which it is useful to express \( L^{\text{latent}} \) as:

\[
L^{\text{latent}}(x, \hat{y}; \phi, \chi) = D_{KL}(q_{\phi}(.|x)||p_{\chi}(.|\hat{y})) \\
= - \int q_{\phi}(z|x) \log \frac{q_{\phi}(z|x)}{p_{\chi}(z|\hat{y})} dz \\
= E_{z \sim q_{\phi}(.|x)} \left[ - \log q_{\phi}(z|x) \right] + E_{z \sim q_{\phi}(.|x)} \left[ \log p_{\chi}(z|\hat{y}) \right] \\
= \frac{1}{2} \sum_{j=1}^{J} \left( 1 + \log(2\pi) + \log \left( \sigma_j(x)^2 \right) \right) + \sum_{c=1}^{N_{\text{num}}} \chi_c N \left( \mu^{(x)} + \sigma^{(x)} \odot \epsilon_c | \mu_c^{e}, \sigma_c^{e2} I \right) \\
\tag{11}
\]

whereby the last equality uses Eq. [9] and the reparameterization \( z = \mu^{(x)} + \sigma^{(x)} \odot \epsilon \). Because of this reparameterization, the Monte Carlo estimate of the final expectation in Eq. [11] is differentiable with respect to \( \phi \). This expectation was again estimated with a single Monte Carlo sample per datapoint (actually using the same sample as for the estimation of the expectation in Eq. [6] or Eq. [12].

When generating a sample to be replayed, the specific class \( y \) to be generated was first randomly selected from the classes seen so far, after which the latent variables \( z \) were sampled from \( N(\mu^{y}, \sigma^{y2} I) \). Although this meant that a specific class was intended to be replayed (and that class could thus be used to label the generated sample with a hard target), it was still the case that the generated inputs were labelled with the (temperature-raised) softmax probabilities obtained for them by a feedforward pass through the network.

**Context gates** To enable context-dependent processing in the generative part of our models, for each class to be learned, a randomly selected subset of \( X\% \) of the units in each hidden layer of the decoder network was fully gated (i.e., their activations were set to zero; Figure [3C]). The value of hyperparameter \( X \) was set by a grid search (Figure [3]). Note that thanks to the combination with conditional replay, during the generation of the samples to be replayed, the specific classes selected to be generated dictated which class-mask to use.

**Internal Replay** To achieve the replay of hidden or internal representations, we removed the de-convolutional or transposed convolutional layers from the decoder network. During reconstruction or generation, samples thus only passed through the fully-connected layers of the decoder. This meant that replayed samples were generated at an intermediate level, and when our model encountered replayed data it let it enter the encoder network after the convolutional layers (Figure [3D]). The reconstruction term of the generative part of the loss function was therefore changed from the...
input level to the hidden level, and it was defined as the expected squared error between the hidden activations of the original input and the corresponding hidden activations after decoding:

\[
L^{\text{recon}}(x; \phi, \psi) = E_{\epsilon \sim \mathcal{N}(0, I)} \left[ \sum_{i=1}^{N_{\text{units}}} \left( h_i(x) - \tilde{h}_i \right)^2 \right] \tag{12}
\]

whereby \( h_i(x) \) was the activation of the \( i \)th hidden unit when the original input image \( x \) was put through the convolutional layers, and \( \tilde{h}_i \) was the activation of the \( i \)th hidden unit after decoding the original input: \( \tilde{h} = p^\ast \psi(z(x)) \) with \( p^\ast \psi \) the decoder network without deconvolutational layers and \( z(x) = \mu(x) + \sigma(x) \odot \epsilon \). The expectation in Eq. 12 was again estimated with a single Monte Carlo sample per datapoint. To prevent large changes to the convolutional layers of the encoder, the convolutional layers were pre-trained on CIFAR-10 (see below) and frozen during the incremental training on CIFAR-100. For a fair comparison, pre-trained convolutional layers were used for all compared methods.

**Other Continual Learning Methods and Baselines Included in Comparison**

**SI / EWC** For these methods a regularization term was added to the loss, with its strength controlled by a hyperparameter: \( L_{\text{total}} = L_{\text{current}} + \lambda L_{\text{regularization}} \). The value of this hyperparameter was set by a grid search (Figure 5). The way the regularization terms of these methods are calculated differs (Zenke et al., 2017; Kirkpatrick et al., 2017), but they both aim to penalize changes to parameters estimated to be important for previously learned tasks.

**LwF** This method (Li & Hoiem, 2017) is very similar to our implementation of standard generative replay, except that instead of using generated inputs for the replay, the inputs of the current task were used. So there was no need to train a generator. As for our version of generative replay, these inputs were replayed after being labeled with soft targets provided by a copy of the model stored after finishing training on the previous task.

**None** As a naive baseline, the base neural network was sequentially trained on all tasks in the standard way. This is also called fine-tuning, and can be seen as a lower bound.

**Joint** To get an upper bound, we sequentially trained the base neural network on all tasks while always using the training data of all tasks so far. This is also referred to as offline training.
Figure 5: Grid searches for EWC, SI, BI-R and BI-R + SI. Shown are the final average test set accuracies (over all 10 tasks / based on all 100 classes) for the hyperparameter-values tested for each method. Note that for the Task-IL scenario, combining BI-R with SI did not result in an improvement, which is why for this scenario the performance of this combination is not reported in the main text. For these grid searches each experiment was run once, after which 10 new runs with different random seeds were executed using the selected hyperparameter-values to obtain the results reported in the main text.