USING THE GNG-M ALGORITHM TO DEAL WITH THE PROBLEM OF CATASTROPHIC FORGETTING IN INCREMENTAL MODELLING

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Abstract: Creating computational models from large and growing datasets is an important issue in current machine learning research, because most modelling approaches can require prohibitive computational resources. This work presents the use of incremental learning algorithms within the framework of an incremental modelling approach. In particular, it presents the GNG-m algorithm, an adaptation of the Growing Neural Gas algorithm (GNG), capable of circumventing the problem of catastrophic forgetting when modelling large datasets in a sequential manner. We illustrate this by comparing the performance of GNG-m with that of the original GNG algorithm, on a vector quantization task. Last but not least, we present the use of GNG-m in an incremental modelling task using a real-world database of temperature, coming from a geographic information system (GIS). The dataset of more than one million multidimensional observations is split in seven parts and then reduced by vector quantization to a codebook of only thousands of prototypes.

1 INTRODUCTION

Building computational models by means of inductive inference has always been an important issue in science. Current information systems gather information in databases containing huge amounts of data, making difficult to tackle the problem of modelling by using traditional methods. Additionally, one must care about the fact that, most of the time, these databases continuously grow, thus requiring a dynamic modelling approach being able to add new knowledge to pre-existent models. As an example, consider modelling a biological process, e.g., growth or development. Biological processes change over time due to the continuous variations on stimuli (e.g., climate). Plants and animals behave different depending on seasons, and seasons evolve through years depending on more complex processes (e.g., global warming). This is the intrinsic complexity of the problem, but this scenario is reinforced by the fact that it is not possible to have all the concerned data in advance. Instead, the data are gradually collected in an incremental way generating growing databases. Creating a model of a such process each time new data are generated could imply large computation time because, in a static modelling framework, the new model must be created from scratch using the whole set of data. A dynamic approach with growing models incorporating the information of new data is desirable in this case.

An interesting approach which deals with this kind of datasets is incremental modelling. This modelling paradigm consists in considering the modelling problem as an incremental task (Giraud-Carrier, 2000), and then using incremental learning techniques in order to build a model from the available data. Therefore, by using this approach, huge datasets may be processed when they are still growing or, they might be transformed by means of a sampling or partitioning procedure¹ before being fed to the incremental modelling framework.

This article presents the use of GNG-m, an adaptation of the popular GNG algorithm (Fritzke, 1995), as a means for achieving incremental learning. More specifically, the idea of the contribution is to use an incremental modelling approach in order to process huge datasets which would require too much memory and computation resources when using standard batch approaches. The remainder of the article contains the following sections. Section 2 introduces the concept of incremental learning, gives some defini-
tions, and shows generalities of the approach. Section 3 explains the standard GNG algorithm, as well as the modifications that were introduced in order to make it more suitable for incremental modelling. Section 4 shows some tests of the performance of the algorithm on a toy-set problem, and Section 5 presents the application of the algorithm to a real-world large multi-dimensional dataset. Finally, Section 6 draws some conclusions about the approach of incremental modelling in the task of processing huge databases, and on the use of incremental learning algorithms like GNG in this purpose.

2 INCREMENTAL MODELLING

The fact of having large or growing datasets, while being positive and also desirable from the point of view of a data acquisition system, constitutes a major drawback if the data are to be used in inductive modelling. In the case of a huge database, one can devise two possibilities. On the one hand, using huge amounts of data to build models could be prohibitive because of computational or storage constraints. And on the other hand, taking only partial information from the complete set and use it to build a model, implies the use of special modelling paradigms being able to append or insert new information into partial or growing adaptive models.

A similar landscape is depicted if one thinks on growing databases. It is a matter of fact that in order to build an accurate model, a large enough amount of data must be available. There are also two difficulties here. On the one hand, it is not a trivial issue to know beforehand the amount of data that is large enough for building a model, and this is even more difficult if the data collecting process is still running. On the other hand, suppose that the amount of gathered information is large enough to build a model. In the case of having a static process as modelling objective, those data would be sufficient to reach some level of accuracy. Conversely, if the model should fit a dynamic process, it would be better to have an adaptive modelling framework that follows the changes of the process, instead of having absolute stable models which have to be rebuilt from scratch each time the process changes. Again, a model structure that incrementally changes with the amount and quality of data might be a good strategy.

A modelling framework which fulfils the aforementioned requirements is incremental modelling, also called incremental learning. In the area of machine learning, the term incremental learning has synonymously been used with pattern learning and online learning to describe the opposite of batch learning (Chalup, 2002). Within this context, it only means to distinguish two policies for modifying the parameters of a model during training i.e., after the presentation of each training example in the online case, or after the cumulation of a certain number of modifications in the batch case. We are considering a more specific definition of the concept of incremental modelling, which will be explained after defining the concept of incremental task in the next subsection.

2.1 Incremental Task

In general, modelling tasks where the examples or observations become available over time (usually one at a time) are considered as incremental learning tasks (Giraud-Carrier, 2000). Traditional static methods can be employed for building a model from an incremental task given that, if there exist the possibility of waiting for the data, any incremental learning task can be transformed into a non-incremental one (Giraud-Carrier, 2000). This approach has the hindrance of reaching excessive volumes of data that could render infeasible the modelling task. It would be preferable to make use of the advantages of incremental learners in this case. In the same way, a large non-incremental modelling task being unbearable by traditional modelling approaches can be transformed into an incremental task by sampling or splitting the data, and then to use an incremental learner in order to build a model from the obtained incremental dataset.

2.2 Incremental Learning

Besides being a synonym of pattern learning or online learning in the machine learning terminology, incremental learning is a concept which has been associated with learning processes where a standard learning mechanism is combined with or is influenced by stepwise adjustments during the learning process. These adaptations can be changes in the structure of the learning system (e.g., growing and constructive neural networks), or changes in its parameters (e.g., stochastic learning), or even changes in the constitution of its input signals (e.g., order, complexity) (Chalup, 2002). These adaptations have the purpose of enabling the construction of more specialized models by adding new information to the already existent knowledge, when it is available.

Within this context, incremental learning shares the same meaning of sequential learning. In sequential learning, the learning system is sequentially trained by using different datasets, which most of the
times are chunks of a larger dataset that grows due to a gathering process which is still running. This definition is very close to the concept of online learning, except but the fact that in pure sequential learning each dataset (and all related information) is discarded after each step of the sequence. Only the model parameters are kept\(^2\) (Sarle, 2002).

Thus, as a conclusion, incremental modelling may be defined as the process of using any incremental or sequential learning algorithm, to solve any incremental learning task, even those which come from the transformation of non-incremental ones. This article emphasizes on the use of incremental learning for solving non-incremental tasks that were transformed into incremental ones by means of a sampling or partition procedure.

### 2.2.1 Stability - Plasticity Dilemma

Incremental learning algorithms must be designed to remain plastic in response to significant new events, yet also remain stable in response to irrelevant events. Moreover, it is desirable that, in the plastic mode of the model, the new events affect only the part of the model being concerned with the new knowledge without causing interference with pre-existent concepts. This compromise between stability and plasticity (Grossberg, 1987; Carpenter and Grossberg, 1987) is difficult to achieve in static models with distributed representations which try to minimize an objective function. This phenomenon is due to the fact that subsequent training datasets may have totally different local minima, making the sequential training of successive datasets to forget all previous sets, resulting in the so called catastrophic interference (McCloskey and Cohen, 1989). Several strategies has been proposed in order to minimize these effects (French, 1994; Robins, 2004) in distributed representations.

Besides a distributed representation, information can also be represented using a local scheme. Local encoding of the information is one of the requirements to successfully perform learning in a sequential manner (Sarle, 2002). The use of models with a local representation of the information allows partial updates of their parameters, permitting new knowledge to be introduced into the model without modifying the previous information already stored. Moreover, local adaptations must be accompanied by the ability of dynamically changing the model structure (i.e., growing), in order to give enough flexibility when required.

\(^2\)In the particular case of artificial neural networks, these parameters are the weighted connections between neurons if knowledge is encoded in a distributed manner, or unit positions if the encoding schema is local.

### 3 THE GROWING NEURAL GAS ALGORITHM

Growing Neural Gas (GNG) (Fritzke, 1995) is an incremental point-based network (Bouchachia et al., 2007) that performs vector quantization and topology learning. The algorithm builds a neural network by incrementally adding units using a competitive Hebbian learning strategy. The resulting structure is a graph of neurons that reproduces the topology of the dataset by keeping the distribution and the dimensionality of the training data (Fritzke, 1997).

The classification performance of GNG is comparable to conventional approaches (Heinke and Hamker, 1998) but has the advantage of being incremental. Hence, giving the possibility of training the network even if the dataset is not completely available all the time while reducing the risk of catastrophic interference by using local encoding.

The algorithm proposed by Fritzke is shown in Table 1. In this algorithm, every \(\lambda\) iterations (step 8) one unit is inserted halfway between the unit \(q\) having the highest error and its neighbour \(f\) having also the highest error. Carrying out this insertion makes the network to converge to a structure where each cell is the prototype for approximately the same number of data points and hence, keeping the original data distribution. The term “nearest unit” in step 2 refers to the more widely used concept of best matching unit (BMU).

As an example, a GNG network was trained using the dataset shown in Figure 1, and the training parameters shown in Table 2. These values were selected after several runs of the algorithm. This dataset contains 10’000 observations drawn from a two-dimensional normal distribution with means \(\mu_x = 5, \mu_y = 5\) and standard deviations \(\sigma_x = 2, \sigma_y = 2\).

Figure 2 shows the position and distribution of

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![Figure 1: Two-dimensional Single-blob normal distribution.](image-url)
Table 1: Original growing neural gas algorithm proposed by Fritze.

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step 0</td>
<td>Start with two units $a$ and $b$ at random positions $w_a$ and $w_b$ in $\mathbb{R}^n$</td>
</tr>
<tr>
<td>Step 1</td>
<td>Generate an input signal $\xi$ according to a (unknown) probability density function $P(\xi)$</td>
</tr>
<tr>
<td>Step 2</td>
<td>Find the nearest unit $s_1$ and the second-nearest unit $s_2$</td>
</tr>
<tr>
<td>Step 3</td>
<td>Increment the age of all edges emanating from $s_1$</td>
</tr>
<tr>
<td>Step 4</td>
<td>Add the squared distance between the input signal and the nearest unit in input space to a local counter variable: $\Delta error(s_1) = |w_{s_1} - \xi|^2$</td>
</tr>
<tr>
<td>Step 5</td>
<td>Move $s_1$ and its direct topological neighbours towards the input signal $\xi$ by fractions $\varepsilon_b$ and $\varepsilon_n$, respectively, of the total distance: $\Delta w_{s_1} = \varepsilon_b (\xi - w_{s_1})$ $\Delta w_n = \varepsilon_n (\xi - w_n)$ for all direct neighbours $n$ of $s_1$</td>
</tr>
<tr>
<td>Step 6</td>
<td>If $s_1$ and $s_2$ are connected by an edge, set the age of this edge to zero. If such an edge does not exist, create it</td>
</tr>
<tr>
<td>Step 7</td>
<td>Remove edges with an age larger than $a_{\text{max}}$. If the remaining units have no emanating edges, remove them as well</td>
</tr>
</tbody>
</table>
| Step 8 | If the number of input signals generated so far is an integer multiple of a parameter $\lambda$, insert a new unit as follows:  
- Determine the unit $q$ with the maximum accumulated error.  
- Insert a new unit $r$ halfway between $q$ and its neighbour $f$ with the largest error variable: $w_r = 0.5 (w_q + w_f)$  
- Insert edges connecting the new unit $r$ with units $q$ and $f$, and remove the original edge between $q$ and $f$.  
- Decrease the error variables of $q$ and $f$ by multiplying them with a constant $\alpha$. Initialize the error variable of $r$ with the new value of the error variable of $q$. |
| Step 9 | Decrease all error variables by multiplying them with a constant $d$ |
| Step 10 | If a stopping criterion (e.g., net size or some performance measure) is not yet fulfilled go to step 1 |

Table 2: Parameters for the Growing Neural Gas algorithm.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>$\varepsilon_b$</th>
<th>$\varepsilon_n$</th>
<th>$\lambda$</th>
<th>$a_{\text{max}}$</th>
<th>$\alpha$</th>
<th>$d$</th>
</tr>
</thead>
<tbody>
<tr>
<td>value</td>
<td>0.005</td>
<td>0.001</td>
<td>500</td>
<td>100</td>
<td>0.5</td>
<td>0.9</td>
</tr>
</tbody>
</table>

Figure 2: Prototype vectors obtained with the GNG algorithm using the whole dataset.

the 50 cells of the resulting structure. A smoothed coloured density representation of the dataset from where the prototypes were obtained was added to every scatter plot in this paper. Showing the original distribution of the training data can be useful to evaluate if the prototypes were correctly placed after the learning process. As we can see in Figure 2, the distribution of each one of the variables is reproduced by the group of prototypes in the network.

Now, in order to simulate the conditions that would require the use of incremental modelling, let us consider this dataset to be so huge that it is impossible to be processed in one single step due to memory constraints. Then, incremental modelling arises as a possible solution to overcome this problem. The huge dataset can be split, transforming the task of modelling into an incremental one, and then one can use an incremental learner (e.g., the GNG algorithm) to build a model of the dataset in a sequential manner.

Figure 3 shows the resulting positions of the pro-
ototypes of the GNG network after doing a sequential training with the split version (2 parts) of the dataset shown in Figure 1. As it can be seen, after the first step of the sequence, the algorithm reproduces the distribution of the first part of the dataset without any problem. Then, after presenting the second part, the algorithm “forgets” the first part (i.e., the quantization error for this part increases), assigning some of the prototypes that represented the first part of the data to the more recent second part of the dataset. This undesirable manifestation of catastrophic forgetting is due to the fact that the algorithm, being pure incremental, does not keep information about previous datasets used to train the network, and therefore, there is no way of knowing if one of the units of the network represented some important data, and that given this fact its position must remain invariable.

In order to minimize the aforementioned situation, one modification to the original algorithm is proposed in Section 3.1.

3.1 The GNG-m Algorithm

The original version of the GNG algorithm does not keep any additional information about the observations that have been presented to the network; only unit positions and edge ages are stored in the model. This policy, while making the algorithm suitable for performing incremental learning (pure sequential algorithm), does not allow the algorithm to know if a unit in the graph has been an important prototype (i.e., a best matching unit) in previous runs.

The idea of the GNG-m algorithm is to attenuate parameters \( e_b \) and \( e_n \) of the original version (which represent the change of position of the units) for the units which have been useful for representing observations previously presented to the network. In order to do that, we associated a property of “mass” to each unit in the network, in a way that units having been useful for quantizing the dataset get more mass than units representing less data. Hence, we adapted step 5 to take into account the mass value of the unit, freeing “lighter” units and locking the “heavier” ones.

In summary, the proposed modification adds two parameters, \( \text{massInc} \) and \( \text{massDec} \), to the original GNG algorithm. Each time a unit is selected as the nearest unit \( s_1 \), its mass is incremented by \( \text{massInc} \), while all its direct neighbours decrease their mass by \( \text{massDec} \). The mass of the unit is then used to modulate how far a unit can move in a single step, and as a result, units having been selected more often as the nearest unit change less their position than units that represent less points in the dataset.

As an example, Figure 4 shows the resulting positions of the prototypes of the GNG-m network after doing a sequential training with the split version (2 parts) of the dataset shown in Figure 1. Contrary to the results shown in Section 3, the modified version of the algorithm satisfactorily places new prototypes when the second part of the file is presented, without modifying the positions of the prototypes representing the first part. More examples using this new algorithm are given in the next section.

4 TOY-SET EXPERIMENTS

This section shows the results of several tests using the dataset shown in Figure 1. Two approaches for modelling a huge dataset are compared i.e., model merging and incremental modelling. In the case of model merging, we explored parallel and cascade merging. For the incremental modelling approach, we used both versions of the GNG algorithm presented in Section 3 and Section 3.1.

In the case of this toy-set, it is possible to train a model with the whole database. Thus, in order to have a reference to compare with, the resulting prototypes when using the whole dataset are shown in Figure 2. The dataset used for the tests has only 10'000 observations; however, we used it in order to simulate a huge one which cannot be loaded in memory. In order
Table 3: Proposed modification to the original algorithm.

<table>
<thead>
<tr>
<th>Step</th>
<th>Move $s_1$ and its direct topological neighbours towards the input signal $\xi$ by fractions $\varepsilon_0$ and $\varepsilon_n$, respectively, of the total distance:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$\Delta w_s = (\varepsilon_p \cdot e^{mass_1}) (\xi - w_s)$ for all direct neighbours $p$ of $s_1$</td>
</tr>
<tr>
<td></td>
<td>$\Delta w_n = (\varepsilon_n \cdot e^{mass_n}) (\xi - w_n)$ for all direct neighbours $n$ of $s_1$</td>
</tr>
<tr>
<td></td>
<td>$mass_1 = mass_{s_1} + massInc$ for all direct neighbours $s$ of $s_1$</td>
</tr>
<tr>
<td></td>
<td>$mass_n = mass_{n} - massDec$ for all direct neighbours $n$ of $s_1$</td>
</tr>
</tbody>
</table>

to do that, this toy-set was split in five and four parts by using different patterns (vertical and tangential respectively).

4.1 Splitting the Dataset in Five Parts

In this first group of experiments the dataset was vertically split in five parts, and each part was used to build a quantized version of the whole set of observations in an incremental manner.

4.1.1 Parallel and Cascade Merging

The first approach we explored for accomplishing the task of building a quantized version of the split dataset is to train small individual models with each part, and then to use the resulting prototypes as inputs for training a final neural network. This final model should reproduce the distribution of the whole dataset.

Figure 5: Prototypes of a GNG after a parallel merging of five sets of pre-computed prototypes.

Figure 5 shows the results of training five models (i.e., one model for each one of the parts of the dataset), and then merging the resulting models into one single final model. This merging is done by taking the positions of all the units in the five neural networks, and using them as inputs to train a final network with the GNG algorithm. As it can be seen, the position of the units of the final model reproduces the input data distribution.

However, this procedure we called parallel merging, only emulates partially our subject of interest. The final model can be trained only after having built each one of the smaller models, or in other words, the whole dataset must be available in order to have a final model reproducing the distribution of the data.

Besides the parallel approach for merging the partial models, we tested a cascade strategy. In this case, a new up-to-date model is incrementally trained each time a new part of the dataset is available. Therefore, a new input dataset must be compiled at each step of the merging strategy by appending the resulting prototypes of the previous step, and the incoming data. This growing dataset is employed for building the model at each step.

Figure 6 shows the results of quantizing the proposed dataset by using a cascade merging approach. As it can be seen in Figure 6(a), the fact of combining a quantized version of the partial dataset (i.e., the positions of the units of the last model) with the incoming data, does not produce a correct representation of the whole dataset. The distribution of units in the GNG algorithm depends strongly on the density of points in the input dataset. The algorithm distributes its units in a way that every unit in the network is a prototype for approximatively the same amount of data points (i.e., entropy minimization). Hence, the algorithm gives more importance to the more populated incoming dataset and tends to “forget” the information of the less populated set of prototypes of the last available model. Figure 6 shows the quantization error after each step of the cascade merging. As it can be seen in Figure 6(b), the quantization error of the first part increases after the presentation of new parts of the dataset.
4.1.2 Incremental Modelling with the GNG and GNG-m Algorithms

In this section we employed the *incremental* approach for building a quantized version of the dataset split in five parts. Figure 7 shows the resulting prototypes when the dataset is sequentially quantized by using the original GNG algorithm. As it was already discussed in section 3, the former parts of the dataset are lost after presenting the last ones because there is no information about how useful each unit is in the network.

Figure 9 shows the quantization error for each part of the dataset after each step of the sequential learning process. As it can be seen in figure 9(a), the quantization error of the former parts increases when the latter parts of the dataset are presented.

As it was mentioned in Section 3.1, the problem of catastrophic forgetting can be solved by using the GNG-m algorithm. Figure 8 shows the resulting prototypes after sequentially training a GNG-m algorithm with the dataset split in five parts. As it
can be seen, the policy of locking units according to its relative relevance (mass of the neurons) makes the algorithm to insert new units where needed, without modifying units which were useful for representing a previous subset.

Figure 9 shows the difference between both versions of the algorithms in terms of the quantization error of each one of the parts of the dataset. As it can be seen in Figure 9(b), the quantization error remains almost constant in the case of the GNG-m algorithm, whereas the GNG algorithm makes it to continuously increase.

**4.2 Splitting the Dataset in Four Parts**

This section shows the results of testing the methodology by using a version of the dataset which was tangentially split in four parts. This splitting pattern allowed us to test how the algorithm behaves when data points close to a former distribution are presented in later stages of the learning sequence. Only results concerning the incremental modelling approach are shown in this section.

Figure 10 shows the positions of the resulting prototypes after sequentially applying the original GNG algorithm to the dataset split in four parts. As it can be seen, the model loses completely the prototypes representing the first part of the dataset after presenting the last one. Conversely, Figure 11 shows the positions of the resulting prototypes after using the GNG-m version of the algorithm. As it can be seen, the modified version of the algorithm manages to keep the information of the first part even after presenting the last part of the dataset. Figure 12 shows the difference in the behaviour of both versions in terms of the quantization error of each part of the dataset, after each step of the sequence.

As it can be seen in Figure 12(b), the quantization error in the GNG-m version of the algorithm remains almost constant, reducing catastrophic forgetting in incremental modelling.

**5 USING A REAL-WORLD DATASET**

In order to further validate the incremental modelling approach, and to give an idea of a real application, we
quantized a real large multidimensional dataset using both versions of the GNG algorithm.

The dataset we used comes from the climate database WORLDCLIM (Hijmans et al., 2005), and contains information of the current conditions of temperature in Colombia (averaged from 1950 to 2000). There are 1’336’025 observations, each one corresponding to one pixel with a spatial resolution of one square kilometre. Each observation has 36 dimensions i.e., maximum temperature for the 12 months of the year, minimum temperature for the 12 months of the year, and average temperature for the 12 months of the year. The goal with this dataset is to build a quantized version with a reduced number of prototypes, which can be analysed faster than taking the whole set of observations. The resulting set of prototypes could be useful for finding homologue regions within the country, or for finding species distributions.

The first step of the process consists in transforming the task of vector quantization into an incremental task. Thus, the dataset was split in 7 parts\textsuperscript{3}, six parts with 200’000 observations, and one last part with 136’025 observations. These parts were used for sequentially training the GNG models. The second step in incremental modelling consists in creating a model from the data by using an incremental learner algorithm. For this application we did not use exactly the original version of the GNG algorithm, but a slightly modified version (Satizabal et al., 2009) which prevents the accumulation of prototypes when the dataset is highly heterogeneous. The parameters we used to quantize the data can be found in Table 4.

The resulting prototypes can not be easily visualised given that the dataset has 36 dimensions. Instead, in order to illustrate the difference between both versions of the algorithm, Figure 13 shows the quantization error of each part of the dataset during all the steps of the sequence of learning. When using the standard GNG algorithm, the error of the first parts increases after presenting the subsequent parts of the dataset. These changes in the quantization error reveal that catastrophic forgetting is occurring during learning. Or in other words, that units which were introduced in the network in order to quantize a specific region of the data distribution, are then reallocated to new regions, causing the quantization error of the former parts to increase.

This situation is alleviated by the modifications proposed in Section 3.1. As it can be seen in Figure 13(b), when the GNG-m version is used, the quantization error of each part of the dataset remains almost constant during the subsequent steps of learning. Instead of moving pre-inserted units, the algorithm inserts new units for the new regions, and keeps the existing ones in order to maintain the knowledge in the network.

6 CONCLUSIONS

Incremental modelling consists in using incremental learning algorithms for building models from data coming from incremental tasks. These incremental tasks can be incremental per se (i.e., where observations are available one at the time), or can be created from non-incremental ones by using partition or sampling procedures. This strategy is useful when one has to deal with large and/or growing databases. In the case of a large database where all the data are available, a sampling or partition procedure can be used to divide the whole dataset into smaller parts. These sub-sets can be used either for building small models which have to be merged into one single final model by using parallel merging, or for feeding an incremental learner which must deal with the construction of an incremental model. Partition and sampling are only possible if the whole dataset is available. In the case of growing databases, the data have to be accumulated to form chunks of data that can be presented sequentially to the incremental learner. In this manner, an up-to-date model is always available as the result of training the last generated model with the more recent data. A merging strategy is not feasible in this case (see Section 4).

In the case of this contribution, we used the Growing Neural Gas (GNG) algorithm as incremental learner to explore the possibility of quantizing a large dataset in an incremental manner. The original algorithm (Fritzke, 1995) was slightly modified by adding two parameters which control the plasticity of the network. In the modified version (GNG-m), each unit in

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\textsuperscript{3}The splitting pattern was only geographic, from north to south, without taking into account the properties of the dataset in the space of features.
Table 4: Parameters for the Growing Neural Gas algorithm.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>$\varepsilon_b$</th>
<th>$\varepsilon_n$</th>
<th>$\lambda$</th>
<th>$a_{max}$</th>
<th>$\alpha$</th>
<th>$d$</th>
<th>hold</th>
<th>$sup$</th>
</tr>
</thead>
<tbody>
<tr>
<td>value</td>
<td>0.05</td>
<td>0.005</td>
<td>250</td>
<td>1000</td>
<td>0.5</td>
<td>0.9</td>
<td>0.1</td>
<td>10%</td>
</tr>
</tbody>
</table>

Figure 13: Average quantization error for each one of the parts at each step.

the network has information of how many times the unit has been selected as the nearest unit of any observation in the dataset. We call this information the mass of the unit. In this way, units having a smaller mass can move more easily than units having a larger mass. The tests we performed showed how these new parameters can be useful in alleviating the problem of catastrophic forgetting in incremental learning.

However, these modifications to the algorithm do not consider the possibility of building models from data coming from dynamic systems. The GNG-m algorithm can increase the property of memory, which represents stability, but at the same time decreases the plasticity of the network. Increasing stability is desired in the case of modelling a large dataset because catastrophic forgetting is undesirable. Conversely, forgetting becomes an important feature when the process generating the data changes over time. In this case, a policy such as “mass-decay” can be a feasible approach to control the ability of the network to forget the oldest events, and thus remaining plastic to capture the changing dynamics of the process being modelled.

REFERENCES


