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To cite this version:
Raphael Couronne, Maxime Louis, Stanley Durrleman. Longitudinal autoencoder for multi-modal disease progression modelling. 2019. hal-02090886v2

HAL Id: hal-02090886
https://hal.archives-ouvertes.fr/hal-02090886v2
Submitted on 17 Apr 2019

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Longitudinal autoencoder for multi-modal disease progression modelling

Raphael Couronne\textsuperscript{1,2}, Maxime Louis\textsuperscript{1,2}, and Stanley Durrleman\textsuperscript{1,2}

\textsuperscript{1} Sorbonne Universites, UPMC Univ Paris 06, Inserm, CNRS, Institut du cerveau et de la moelle (ICM)
\textsuperscript{2} Inria Paris, Aramis project-team, 75013, Paris, France

Abstract. Imaging modalities and clinical measurement, as well as their time progression can be seen as heterogeneous observations of the same underlying disease process. The analysis of sequences of multi-modal observations, where not all modalities are present at each visit, is a challenging task. In this paper, we propose a multi-modal autoencoder for longitudinal data. The sequences of observations for each modality are encoded using a recurrent network into a latent variable. The variables for the different modalities are then fused into a common variable which describes a linear trajectory in a low-dimensional latent space. This latent space is mapped into the multi-modal observation space using separate decoders for each modality. We first illustrate the stability of the proposed model through simple scalar experiments. Then, we illustrate how information can be conveyed from one modality to refine predictions about the future using the learned autoencoder. Finally, we apply this approach to the prediction of future MRI for Alzheimer’s patients.

Keywords: Longitudinal · Multi-modal · Autoencoder

1 Introduction

The longitudinal pattern of progression of a disease contains more information than a static observation. Leveraging this information is a key problem in machine learning for healthcare, complicated by to the nature of clinical datasets. These datasets may contain very heterogeneous observations from various modalities of subjects at multiple time points, such as clinical scores, imaging and biological samples. They include missing values, often by design: not all modalities are observed at each visit. Besides, the number of observations and their time spacing vary between subjects. For these reasons, the analysis of multiple modalities and their time dynamic at once is a challenging task.

Linear mixed effect model estimated via EM and their extension to the non-linear case \cite{4,5} were developed for the analysis of unimodal longitudinal data. More recently, recurrent auto-encoder \cite{9,11} offer a way to encode trajectories into a low-dimensional embedding, allowing to perform unsupervised clustering of the trajectories \cite{2}. Riemannian geometry based approaches such as \cite{6,10} offer ways to learn sub-manifolds of the observation space with a system of coordinate adapted to the progression of the modality observed in the data.
On the other hand, various unsupervised methods exist to fuse information from multiple modalities but from a single time snapshot. In [1, 8], the authors propose to learn a common embedding for multiple modalities auto-encoding, merging the information from all modalities and allowing the generation of missing modalities. In [7], unsupervised features are learned from heterogeneous health data as a dimensionality reduction method before machine learning tasks. In [12], combining time and multi-modal approaches, the authors propose a setting for multi-modal time-series embedding. But their design does not handle missing modalities, common in clinical data sets. Besides, the fusion of the information from the different modalities is done at each time step and not on the progression pattern globally, thus decreasing the importance of the dynamics of each modality in the encoding.

To address these limitations, we propose a new setting for longitudinal multi-modal encoding. We extend to the multi-modal case the approach of [6]. Each modality is first separately encoded using a recurrent neural network. A fusion network is then used to merge the obtained representations into a unique representation, which describes the multi-modal trajectory of the subject as a time-parametrized linear trajectory in a latent space $Z$. Then, this trajectory is decoded using a different neural network for each modality, which generates continuously varying trajectories of data changes. This setting allows to handle multiple modalities even when not all of them are observed at each visit and it can handle any number of visits and any time spacing between the visits. Finally, extrapolation in the latent space allows for prediction of the future of each modality and we show on a synthetic dataset and on the ADNI database using cognitive scores and MRI jointly that the predictive power is enhanced by the fusion of each modality embeddings.

In section 2 we explain the proposed model, in section 3 we present experimental results highlighting the stability of the method on synthetic and real data sets and we show how the information from one modality that contributes to the encoding allows to refine prediction of the future of another modality.

2 Methods

We set a longitudinal dataset which contains repeated observations of subjects, where the observations at each time point contain a various combination of modalities among $M \in \mathbb{N}$ modalities. For any subject $i \in \{1, \ldots, N\}$ where $N \in \mathbb{N}$ and for any modality $m \in \{1, \ldots, M\}$, we have a sequence $(y_{ij}^m, t_{ij}^m)_{j=1,\ldots,n_m}$ of observations $y_{ij}^m$ of observed at times $t_{ij}^m$.

2.1 Decoding : Non linear mixed effect model

We set $d \in \mathbb{N}$ and consider a $d$-dimensional latent space $Z = \mathbb{R}^d$ and its canonical basis $(e_i)_{i=1,\ldots,d}$. Then, in the spirit of random slopes and intercepts models, we consider trajectories in $Z$ of the form $l(t) = e^\eta (t-\tau) e_1 + \sum_{i=2}^d \lambda_i e_i$ where $\eta, \tau, \lambda_2, \ldots, \lambda_d \in \mathbb{R}$ are random variables. These trajectories progress in the $e_1$
individual parameters $\phi_i$ are estimated via the use of an encoder network. More precisely, each modality is first processed by a dedicated Recurrent Neural Network (RNN), to get modality-wise representations. To correct for the varying spacings between the observations, we provide to the RNN the visit times, previously normalized to zero-mean and unit variance.

We then concatenate the obtained representations, and use a fully-connected network to merge the representations. The given architecture allows fast inference for new subjects, and is trainable end to end. Besides, the fusion operation is learned so as to produce a single vector which contains the most information about the reconstruction of the whole sequences of all the modalities. The left part of Figure 1 illustrates the procedure.

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**Fig. 1:** Description of the proposed longitudinal autoencoder.
2.3 Regularization, cost function and optimization

To enforce some structure in the latent space and in the family of trajectories obtained, we set the following regularization on the individual variable $\Phi_i$:

$$r(\eta, \tau, (\lambda_i)^{i=2,\ldots,d}) = \eta^2 + \tau^2 + \sum_{j=2}^{d}(\lambda_j)^2.$$  

This regularization models the $\eta$ variable to be distributed along a zero-centered normal distribution, which allows the pace of progression to vary typically between 0.2 and 5 times the mean velocity. The $\tau$ variable is regularized the same way. This regularization is not arbitrary: during each run, the observation times $t_{ij}$ are rescaled to zero-mean unit variance, and thus $\tau$ can handle delays between subjects of order the standard deviation of the observation ages.

Overall, the optimized cost function for one subject is the regularization cost added to the $\ell^2$ reconstruction cost summed over all modalities:

$$C((w_m)_m, \eta, \tau, (\lambda_i)_i) = r(\eta, \tau, (\lambda_i)_i) + \sum_m \frac{1}{\sigma^2_m} \sum_{j=1}^{n_{ij}} \|y_{ij}^m - \Psi_w(l_i(t_{ij}^m))\|^2_2$$ (1)

where the $(\sigma^2_m)_m$ are trade-off parameters between each modality and the regularization. We set an automatic update rule for these parameters after each batch by setting them to the empirical quadratic errors in reconstruction for the modality over the batch. The estimation is achieved by stochastic gradient descent with the Adam optimizer [3] and a batch size of 32 subjects. The Decoders are either fully connected or de-convolution networks depending on the kind of modality considered, with standard architectures. The Decoders are either Elman networks or Elman networks working on features extracted using a convolution network in the case of images. All networks are trained end to end using back-propagation and the PyTorch library. A complete code to reproduce these experiments will be released upon publication of the paper.

3 Experimental results

3.1 Cognitive scores: proof of concept

As in [10], we apply our model on repeated measurement of 4 normalized cognitive score extracted from the ADNI cohort, respectively associated with memory, language, praxis and concentration. We include the 248 MCI-converter subjects, followed for an average of 3 years, over 6 visits. We conduct 2 experiments in order to assess the robustness of the method, and report estimated average trajectories in Figure [2] as well as individual reconstruction errors in Table [1] computed from a patient-wise 10-fold cross validation.

First, we apply our model on an increasing partitioning of input feature. We consider 3 cases: selecting all scores at once as one modality, selecting separately memory+language and praxis+concentration as two modalities, and selecting each one separately. We note the overall good stability of the average model over multiple multi-modal architectures, with stability decreasing in the 4-modalities scenario, arguing for a concatenation of the consistent features.
In our second experiment, we assess the robustness of the model with the number of visits per subjects. To this end we consider the 2-modalities scenario, and perform a pruning of the dataset, removing an increasing number of visits of the second modality, i.e. praxis+concentration per subjects. Datasets are obtained from pruning frequencies of respectively 10%, 20% and 40%. Here we also observe an overall good stability of the average trajectory over pruning frequency.

### 3.2 A synthetic dataset

To test the proposed setup in realistic conditions, we generate a synthetic multi-modal data set comprising 300 subjects observed 7 times on average. The first modality is a 2D image of a cross, with varying arm lengths and angles while the second modality consists of two scores with a sigmoid-like growth. We set a time reparametrization function \( s \) with parameters \( a_1, a_2 \) defined by:

\[
s_{a,b}(t) = \frac{1}{1 + e^{-a_2(t-a_1)}}
\]

<table>
<thead>
<tr>
<th>Partitioning</th>
<th>Pruning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-mod</td>
<td>2-mod</td>
</tr>
<tr>
<td>Train ( (x10^{-3}) )</td>
<td>6.7 / 3.8 / 9.7</td>
</tr>
<tr>
<td>Test ( (x10^{-3}) )</td>
<td>7.8 / 5.1 / 10.6</td>
</tr>
</tbody>
</table>

Table 1: Mean 10-fold reconstruction error for the 2 cognitive scores experiments for each modality respectively
Fig. 3: Left: average trajectory and reconstruction examples for the scalar data. Right: average trajectory and some reconstructions for the image data.

t + \text{sign}(t)t^2 + bt^3. To generate an individual, we sample two sets of parameters \((a_k, b_k)_{k=1,2}\). These serve to reparametrize a scenario of score increase: the \(k\)-th score for the subject at time \(t\) is given by \(\sigma \circ s_{a_k, b_k}\) where \(\sigma\) is the sigmoid function. Then, the arms lengths \(L_1, L_2\) for the images of the subject at time \(t\) are given by \(L_1 = \sigma \circ s_{a_2-a_1} + \epsilon_{a_1}, (b_2-b_1) + \epsilon_{b_1}\), \(L_2 = \sigma \circ s_{a_2+a_1} + \epsilon_{a_2}, (b_2+b_1) + \epsilon_{b_2}\) where the \(\epsilon\) are samples from a zero-mean normal distribution and constant with time. Finally, the arm angles are sampled along a normal distribution but are not informative of the synthetic disease process. This design is so that the images contain, in an intricate way, information about the progression of the scores materialized through the \(a_1, a_2, b_1, b_2\) variables. The two modalities are different noisy facets of a common underlying process.

We perform a patient-wise 10-fold estimation of the model this data set. Figure 3 shows the obtained average trajectory for the first fold, as well as the reconstructions of some subjects images and scores observations. We evaluate and average for all folds the test and train reconstruction errors. For the cross, the test error is \(2.0 \times 10^{-8} \pm 8 \times 10^{-9}\) while the train error is \(1.7 \times 10^{-8} \pm 3.9 \times 10^{-9}\). For the scores, the test error is \(7.10^{-3} \pm 3.10^{-3}\) while the train error is \(7.10^{-3} \pm 3.10^{-3}\). This shows that the model generalizes well to unseen data.

We use the trained model to predict the future scores on the test data. We do so by decoding the extrapolation of the latent trajectory encoded by the model. We repeat this experiment by gradually removing the last observations of the image modality, to look at the impact of this modality on the predictive power of the model. Figure 4 shows the experimental setup and the results. As the time span of the observed images shrinks, the prediction deteriorates: when more image data is available, the score prediction is more accurate. This shows the ability of the model to find a relevant common representation for the progressions of the different modalities.
3.3 Application to Alzheimer’s disease future image prediction

On the 248 patients of section 3.1, we apply the same model on the 217 that have at least 1 MRI observation, leading to a total of 1199 cognitive scores measurements and 1441 MRIs. We work on both the MRI images and the cognitive scores. The MRI images are rigidly aligned and sub-sampled to $64^3$ resolution. Note that the subjects do not have both the MRI and the cognitive scores measurements at each visit.

Figure 5 shows one of the estimated average trajectory for the MRI modality. We evaluate and average for all folds the test and train reconstruction errors on both modalities. For the MRI, the test error is $2.5 \times 10^{-3} \pm 6 \times 10^{-5}$ while the train error is $2.4 \times 10^{-3} \pm 2 \times 10^{-5}$. For the scores, the test error is $2.2 \times 10^{-2} \pm 3 \times 10^{-3}$ while the train error is $1.7 \times 10^{-3} \pm 6 \times 10^{-4}$. This shows that the model generalizes well to unseen data.

We then perform the same prediction task as in the previous section: we attempt to predict the future MRI from past data, using a variable amount of score data in the past. Figure 5 shows the prediction errors for different time horizon. Once again, the errors increase as we feed the model with less cognitive scores measurements. This shows that the model captures information contained in the cognitive scores progression to refine the MRI prediction.

4 Conclusion and perspectives

We extended on a deep autoencoder architecture with a mixed effect latent space to propose a practical framework for modeling multi-modal longitudinal data, trainable end-to-end. This allows for analysis of heterogeneous longitudinal datasets, deriving a model-wise average trajectory, as well as condensed patient representations. We study its robustness toward modalities partitioning and

Fig. 4: Left: description of the prediction setup. Right: the MRI prediction errors.
dataset pruning and illustrate its utility in both synthetic and real scenarios. In the future we plan to model the progression of more modalities at once. This work has been partially funded by the European Research Council (ERC) under grant agreement No 678304, European Unions Horizon 2020 research and innovation programme under grant agreement No 666992, and the program Investissements davenir ANR-10-IAIHU-06.

References