Unsupervised Representation Learning of Dynamic Retinal Image Changes by Predicting the Follow-up Image

Antoine Rivail*  Hrvoje Bogunović  Sebastian M. Waldstein
Bianca S. Gerendas  Wolf-Dieter Vogl  Ursula Schmidt-Erfurth
Christian Doppler Laboratory for Ophthalmic Image Analysis
Department of Ophthalmology and Optometry
Medical University of Vienna

1 Introduction

Longitudinal imaging allows to capture both, the static anatomical structures and the dynamic changes of the morphology due to aging or disease progression. However, common supervised or unsupervised methods for medical imaging do not consider dynamic aspects and process longitudinal data as individual data points. For natural images, algorithms already exist that learn a representation from videos [Walker et al., 2016]. In retinal imaging, however, the temporal sampling resulting from follow-up to disease progression is much lower than in videos. Predictions are therefore more ambiguous and prone to noise.

We propose a deep learning approach to overcome these challenges, which allows us to understand the underlying morphological organization and its changes over time, and to discover abnormalities and pathologic evolutions. Our data-driven approach learns a feature representation from unlabeled longitudinal images by predicting the unobserved subsequent image within a series of observations. Several sources of noise, such as imaging noise, misalignment of follow-up images or motion artifacts aggravates the direct prediction of the target image. Thus, we propose to adapt a Conditional Variational Autoencoder (CVAE) [Kingma and Welling] to learn representative static and dynamic features that are robust to noise and uncertainty.

2 Method

Theory of conditional variational autoencoders

Let $X_i = [x_{i-j}, \ldots, x_i]$ be a sequence of $J$ consecutive images $x_i$, from which a subsequent unseen image $x_{i+1}$ is predicted. Following [Walker et al., 2016] [Kingma and Welling], the optimization of variational autoencoder is based on the variational inequality, which maximizes the likelihood of the prediction $P(x_{i+1}|X_i)$ by optimizing the last term of the equation 1. $Q$ is the introduced distribution.

$$
\log P(x_{i+1}|X_i) \geq \log P(x_{i+1}|X_i) - \mathcal{KL}[Q(z|X_i, x_{i+1})|| P(z|X_i, x_{i+1})]
$$

$$
= E_{z \sim Q}[\log P(x_{i+1}|z, X_i)] - \mathcal{KL}[Q(z|X_i, x_{i+1})|| P(z|X_i)]
$$

The term $E_{z \sim Q}[\log P(x_{i+1}|z, X_i)]$ encourages the model to output a correct prediction, where $X_i$ is encoded by the Encoder network. This term is optimized with a $L_2$ reconstruction loss. The second term, the $\mathcal{KL}$ divergence, forces $Q$ to be a normal distribution ($\mathcal{N}(0, I)$). Thus, the encoding network is forced to extract as much information as possible from the previous images. At testing time, the $Q$ network is discarded, and $z \sim \mathcal{N}(0, I)$.

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Figure 1: Architecture: the encoder, the $Q$ Network ($\mu$ and $\sigma$) and decoder are implemented by convolutional networks. At training time $z = \mu + \epsilon \cdot \sigma$, at testing time: $z \sim N(0, I)$

Figure 2: Example of four subsequent OCT B-scans acquired in a monthly interval. The first three images are used as input and the last one as target (red). Retinal thickness, hyperreflective foci or Drusens are morphological properties tested for feature evaluation.

CVAE is computing a distribution over the possible solutions instead of a single prediction as for instance an autoencoder provides. This distribution is built from two parameters: the code vector, $h$, which encodes deterministic factors and the $z$ vector, which represents unpredictable factors.

Architecture: the optimization relies on three convolutional networks: (1) the encoder network, which extracts important information for predictions in the code vector $h$, (2) the $Q$ network for implementing a standard distribution $N(\mu(X_i, x_{i+1}), \sigma(X_i, x_{i+1}))$, and (3) the decoder having as inputs the vectors $h$ and $z$ and which outputs a prediction of the subsequent image (Figure 1).

3 Experiments and results

3.1 Dataset

The dataset contains 3900 OCT scans from 204 different patients diagnosed with intermediate age-related macular degeneration (AMD). Each patient was scanned with a monthly follow-up for a period of up to 24 months. Time-points where a patient already converted to late stage AMD were excluded.

Preprocessing: We built sequences of four consecutive visits without overlapping, and cropped local patches from the central 3 mm of the retina. In order to crop patches from the same anatomical position, we registered all scans to a reference scan within the sequence as described by Vogl et al. [2017]. Finally, every sample is a sequence of four 170 x 170 patches. The first three images were used as inputs to predict the last one (Figure 2).

3.2 Training

We divided the dataset into training and validation subsets for the training of the CVAE and a test set for the final evaluations. Care has been taken that all image series of a patient were in the same subset. The network was trained by stochastic gradient descent using ADAM algorithm. Overfitting was controlled by examining the reconstruction loss on the test set (without Q network).
Table 1: Prediction results of morphological properties. “Direct” shows the results predicted from previous values of the properties only. “CVAE” uses in addition the code vector $h$ from previous images. The target values are scaled to have mean and unit variance (standard scaling).

<table>
<thead>
<tr>
<th>Method</th>
<th>Retinal thickness</th>
<th>Drusen Volume</th>
<th>HRF volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>R² score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Direct</td>
<td>0.959</td>
<td>0.945</td>
<td>0.807</td>
</tr>
<tr>
<td>CVAE</td>
<td></td>
<td>0.974</td>
<td>0.823</td>
</tr>
<tr>
<td>Mean absolute error (standard scaled)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Direct</td>
<td>0.152</td>
<td>0.228</td>
<td>0.406</td>
</tr>
<tr>
<td>CVAE</td>
<td>0.177</td>
<td>0.185</td>
<td>0.392</td>
</tr>
</tbody>
</table>

3.3 Evaluation

In order to evaluate the features produced by the encoder ($h$ code), we predicted morphological properties from it, which change over time in intermediate stage of AMD. We used average total retinal thickness, drusen volume and hyperreflective foci (HRF) volume, which were automatically segmented using the methods described in [Garvin et al., 2009, Schlegl et al., 2017].

The prediction target was the average value of a morphological property from the last image in the sequence (unobserved image). We trained a self-normalizing MLP regressor [Klambauer et al., 2017]. For comparison, we performed a regression based on the measured property values from the initial time-points serving as baseline (Direct). In the CVAE method, we in addition included the code vector $h$ as regression input. Results are listed in Table 1.

Initial results showed that the prediction code $h$ improves the results of a direct prediction, both for Drusen Volume and HRF volume, by increasing R² score and reducing the mean absolute error. This indicates that the proposed method is able to successfully encode dynamic properties of OCT images. The prediction of total retinal thickness was not improved but given that in intermediate AMD total retinal thickness usually remains very stable, it can be predicted more easily by simply regressing from previous thickness measurements.

References


