Can liquid crystal phases be identified via machine learning?

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Abstract

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1 Introduction

Machine learning methods have seen widespread utilisation across all scientific disciplines, in situations where conventional algorithms are too cumbersome to implement for specific data-based and modelling tasks [1]. Deep learning, loosely defined as machine learning with large datasets, parallel computation and scalable algorithms with many layers [2], has and continues to increase the range and complexity of possible applications of machine learning in the sciences [1]. Any task applying deep learning to data with a grid-like form, such as images, likely involves the usage of convolutional neural network (CNN) algorithms [2]. CNNs were conceived in 1989 by Yann LeCun et al. and successfully applied to recognition of handwritten characters [3]. However, their astounding performance in the field of computer vision would not be fully realised until after breakthroughs in deep learning starting in 2006 [2]. Their efficacy was further proven when Geoffrey Hinton et al. entered a CNN into the ImageNet Large Scale Visual Recognition Challenge in 2012, and won by a large margin [4].

Liquid crystal phases are in general identified by eye, directly from textures taken by polarised microscopy. Without adequate experience, this can prove a difficult task because certain unique liquid crystal phases, generated by often minor changes in structural properties, can have similar textural appearances [5]. Our project aims to test the viability of machine learning algorithms as tools to assist phase identification. CNNs are particularly suitable due to their prevalence in image classification, and so form the core of our investigations. Current literature in this specific topic is limited, and the approaches so far have mostly involved the usage of simulated textures in the training of models [6, 7]. Sigaki et al. have demonstrated the viability of CNNs in isotropic and nematic phase texture classification and in the prediction of physical liquid crystal properties [6]. Our study further explores and attempts to push the limits of the classification task across a wider range of phases, utilising real experimental data produced by polarised microscopy.

This project report will first provide a brief overview of the physics behind liquid crystals and the capturing of their textures by polarised microscopy, as well as an introduction to machine learning, neural networks and CNNs. The details and results of our investigations into phase classification will then be presented, as well as an outlook to further study.

2 Liquid crystal phases

Liquid crystals are substances in a state between that of a fully isotropic liquid and a crystal with a periodic lattice structure. The molecules can have varying positional order, and have orientational order over large sections. The unit vector parallel to

the alignment of the molecules is called the director. Other details such as molecular shape and chirality affect the overall structure. These variations in structure result in numerous individual identifiable liquid crystal phases. Thermotropic liquid crystals exhibit phases transitions with changing temperature, whereas lyotropic liquid crystals are dissolved in a solvent with the phase depending on the concentration [8]. This project will be concerned with only thermotropic liquid crystals.

When cooling a thermotropic liquid crystal starting as an isotropic liquid, it will first transition to the nematic phase (N), which has orientational order only. The chiral nematic (cholesteric, N*), phase also has no positional order, and has a periodic variation of the director, resulting in helical structures. Upon further cooling, the smectic phase will be reached. This can be split into three categories, going from fluid smectic to hexatic smectic to soft crystal in order of decreasing temperature. The fluid smectic phase has molecules arranged in layers, with no positional order in the plane of each layer. When the director is perpendicular to the layer planes, the phase is smectic A (SmA), with smectic C (SmC) having a director that is tilted by comparison. Hexatic smectic phases have short range positional order within the layer planes with hexagonal intermolecular arrangements interspersed with orderbreaking defects. This encompasses the smectic B (SmB), I (SmI), and F (SmF) phases. Semctic B has a director perpendicular to the layer planes, whereas it is tilted towards the vertices of the hexagons for smectic I and towards to the sides of the hexagons for smectic F. The soft crystal phases are defect free within the layers and therefore exhibit long range positional order [5].

The liquid crystal texture data used in this project have all been obtained by polarised microscopy captured with a video camera. In brief terms, a polarising microscope works by placing a sample between perpendicularly aligned polarisers. When light is shone through the arrangement the resulting image will be dark, unless the sample rotates the plane of polarisation. In the case of liquid crystals, the isotropic liquid phase has no optical properties so will produce completely dark textures. The nematic, cholesteric and smectic phases are anisotropic and therefore birefringent, with optical axes depending on their structures. This produces unique textural image features for each phase [5]. Some example textures taken from our dataset are displayed in Figure 1.

3 General machine learning principles

A machine learning model is a computer algorithm which automatically improves its performance in a given task as it gains experience from a dataset [2]. It learns patterns in data and uses these patterns to make probabilistic predictions. The data normally takes the form of a set of N examples, which are usually expressed as vectors, or

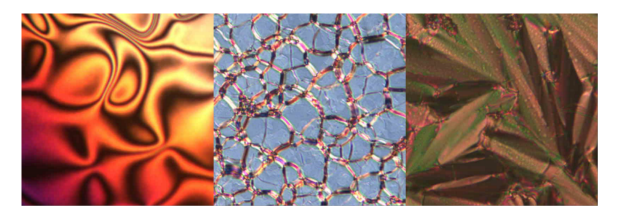


Figure 1: Liquid crystal textures from the dataset, from left to right: nematic phase compound 5CB, cholesteric phase compound D5, and smectic C phase compound M10.

some other structure of features, $\{x^{(i)}\}_{i=1}^N$, containing quantitative information about example i. In supervised learning the examples are given labels $y^{(i)}$, to form a training set of pairs $\{x^{(i)}, y^{(i)}\}_{i=1}^N$, and the model attempts to learn the mapping from a general input x to an output \hat{y} . One main type of supervised learning is regression, in which the output is a numerical scalar. The other main type is classification, in which the model predicts what the input belongs to out of a selection of classes. In unsupervised learning there are no labels, and the algorithm attempts to learn specific patterns in the dataset such as clusters of similar data points [9]. The topic of this project is a supervised classification problem.

A supervised model can be usually be expressed as a function of inputs and a set of parameters $\boldsymbol{\theta}$ such that $\hat{y} = f(\boldsymbol{x}; \boldsymbol{\theta})$. Training involves optimisation of the parameters by minimisation of a cost function $J(\boldsymbol{\theta})$, which measures the deviation of the predictions of the model from the true labels. The most common optimisation algorithms involve computing the gradient of $J(\boldsymbol{\theta})$ with respect to $\boldsymbol{\theta}$ [2].

The capacity of a model is akin to its complexity. The number of trainable parameters can give a fast indication of capacity. However, it also depends on the model's functional form. The parameters controlling the capacity of the model, as well as certain other training settings, are known as hyperparameters. If the capacity is too small, the model will tend to "underfit" the training set, resulting in poor performance even when optimised well. On the other hand, too high a capacity will result in "overfitting", with high performance on the training set, but the model may not generalise well to new unseen data. Methods for reducing overfitting are known as regularisation. Before training begins, the entire dataset is often split into training, validation, and test sets, containing $N_{\rm train}$, $N_{\rm valid}$ and $N_{\rm test}$ examples respectively. The

training set, as defined previously, is used to optimise the parameters. The model's performance is then evaluated on the validation set. A poor performance on both the training and validation sets is indicative of underfitting, whereas a high performance on the training set and low on the validation set suggests overfitting. The validation set can therefore be used to tune the hyperparameters of the model before retraining. This can be repeated until the model fits optimally. The final model is then evaluated on the as-of-yet unseen test set to determine its overall efficacy [2].

4 Feedforward neural networks

A neural network is a type of machine learning model that takes inspiration from the current understanding of how the brain works. The inputs are "fed forward" through a series of connected "hidden units" that represent individual neurons in a brain, before reaching the output units. In the most basic form, a fully connected neural network, The units are arranged into layers, with each unit in a layer connected to every unit of the previous layer. We will define the total number of layers, excluding the input layer, as the depth, D, of the model, with the width, W_l , of layer $l \in [0..D]$ equal to the number of units it contains. l = 0 is the input layer. The choice of the hyperparameters D and W_l defines the architecture of the model. A schematic of an example network is presented in Figure 2.

A single hidden unit's output value is calculated by multiplying the output of each of the previous layer's units with a weight parameter, summing these together with a bias parameter, and then passing the result through a non-linear activation function. More formally, the output O_{lu} of hidden unit $u \in [1..W_l]$, of layer l, with bias parameter b_{lu} and activation function A(h) is calculated as

$$O_{lu} = A(h_{lu}) \tag{1}$$

with

$$h_{lu} = b_{lu} + \sum_{v=1}^{W_{l-1}} \theta_{luv} O_{(l-1)v}$$
 (2)

for l > 0. θ_{luv} is the weight parameter that hidden unit u of layer l applies to the output of unit $v \in [1..W_{l-1}]$ of the previous layer. O_{0u} , is equivalent to value x_u of the the input vector \boldsymbol{x} . In matrix form,

$$O_l = A(h_l) \tag{3}$$

with

$$\boldsymbol{h}_l = \boldsymbol{\theta}_l \boldsymbol{O}_{l-1} + \boldsymbol{b}_l \tag{4}$$

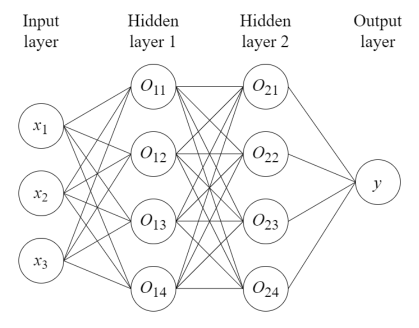


Figure 2: Diagram of a feedforward fully-connected neural network with three input values, D = 3, $W_1 = W_2 = 4$ and one output unit.

where $\boldsymbol{\theta}_l$ is the $W_l \times (W_{l-1})$ dimensional matrix of weights for layer l, with rows corresponding to the weights of each hidden unit, \boldsymbol{O}_l is the vector of outputs for layer l, and \boldsymbol{b}_l is the vector of bias parameters for layer l. The activation function is applied element-wise. For the first layer,

$$\boldsymbol{h}_1 = \boldsymbol{\theta}_1 \boldsymbol{x} + \boldsymbol{b}_1. \tag{5}$$

The number of units in the final output layer depends on the type of model. For regression, there will be one unit that outputs a continuous-valued prediction. For classification, the number of final layer units is equal to the number of classes, $C = W_D$, with each one outputting the predicted probability that an input \boldsymbol{x} belongs to a particular class. The network is therefore summarised as $\hat{\boldsymbol{y}} = f(\boldsymbol{x}; \boldsymbol{\theta})$, with $\hat{\boldsymbol{y}}$ being the C dimensional vector of output probabilities. Each example in the dataset is labelled by a vector \boldsymbol{y} with a value of one for the component corresponding to the true class, and zero for all other components. The component y_u , with $u \in [1..C]$, is equivalent to the Kronecker delta δ_{ut} where the index t corresponds to the true class. The most common choice of final layer activation function for classification is the softmax function, $\sigma_{SM}(h)$, in which case the components \hat{y}_u of the output probability prediction vector, are given by

$$\hat{y}_u = O_{Du} = \frac{e^{h_{Du}}}{\sum_{v=1}^C e^{h_{Dv}}}.$$
(6)

When there are just two classes, known as binary classification, only one unit is needed in the output layer. The data labels y are equal to one or zero depending on the class that x belongs to. The activation function in this case is generally the logistic sigmoid function, $\sigma(h)$, with

$$\hat{y} = O_D = \frac{1}{1 + e^{-h_D}} \tag{7}$$

where in this situation $\boldsymbol{\theta}_D$ is a W_{D-1} dimensional vector. The value of \hat{y} is the predicted probability that \boldsymbol{x} belongs to one of the classes, with a probability of $1-\hat{y}$ that it belongs to the other. The current most commonly used activation function for hidden units is the rectified linear unit (ReLU),

$$A(h) = \max(0, h). \tag{8}$$

In 1989 Kurt Hornik *et al.* mathematically proved that feedforward neural networks with multiple layers and non-linear activations can approximate any continuous function given the correct configuration [10].

5 Neural network training

Neural network training starts with random initialisation of the weights, for example by drawing values from a normal distribution. A single update step is generally carried out by calculating the model output, followed by the derivative of the cost function with respect to the model weight parameters, for each example in the training set. An optimisation algorithm then uses the gradients to update the parameters. Second order optimisation algorithms also utilise the second derivative of the cost function. Generally, the cost function is the expectation value of the loss of all examples in the training set,

$$J(\boldsymbol{\theta}) = \frac{1}{N_{\text{train}}} \sum_{i=1}^{N_{\text{train}}} L(\boldsymbol{f}(\boldsymbol{x}^{(i)}; \boldsymbol{\theta}), \boldsymbol{y}^{(i)})$$
(9)

where the loss is the cross-entropy between the model's output probabilities and the input's true class label,

$$L(\boldsymbol{f}(\boldsymbol{x};\boldsymbol{\theta}),\boldsymbol{y}) = L(\hat{\boldsymbol{y}},\boldsymbol{y}) = -\sum_{u=1}^{C} y_u \log(\hat{y}_u).$$
(10)

When training a classifier model, the softmax activation function of the final layer can be included in the loss function, to give the categorical cross-entropy,

$$L_{CCE}(\hat{\boldsymbol{y}}, \boldsymbol{y}) = -\sum_{u=1}^{C} y_u \log \sigma_{SM}(h_{Du}) = -\log \left(\frac{e^{h_{Dt}}}{\sum_{j=1}^{C} e^{h_{Dj}}} \right)$$
(11)

where we have used the fact that $y_u = \delta_{ut}$ for classification. For binary classification in which the logistic sigmoid activation function is used, the binary cross-entropy is

$$L_{BCE}(\hat{y}, y) = \begin{cases} -\log \sigma(h_D) & \text{if } y^{\text{(true)}} = 1\\ -\log (1 - \sigma(h_D)) & \text{if } y^{\text{(true)}} = 0 \end{cases}$$
 (12)

6 Neural network regularisation

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7 Convolutional neural networks

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7.1 Convolutional layers

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7.2 Pooling layers

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8 4-phase classifier models

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8.1 Dataset preparation

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8.2 Model architectures and training configuration

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8.3 Results

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