1. Introduction

Motivation and goals

The recognition of shape characteristics is one of the major aspects of visual information processing. Together with colour, motion and depth processing it forms the main pathways in the visual cortex. Experiments in cognitive science show in a quite impressive way, that humans recognise complex shapes by decomposition into simpler parts and interpreting the former as coherent spatial compositions of these parts [7]. Corresponding guiding principles for the decomposition are identified from these experiments as well as from research in computer vision (see e.g. [9]). The formulation of these principles relies however on the assumption that the objects are already segmented and thus concepts like convexity and curvature can be applied.

From the point of view of computer vision it is desirable to use shape processing and modeling in the early stages of visual processing. This allows to control e.g. segmentation directly by prior assumptions or by feedback from higher processing layers. This leads to the question whether compositive shape models can be represented and learned in a topologically fully distributed way. The aim of the presented work is to study this question for probabilistic graphical models.

Related work

All mathematically well principled shape models for early vision can be roughly divided into the following two groups.

Global models treat shapes as a whole. Prominent representatives are variational models and level set methods in particular. A shape is described up to its pose by means of a level set function defined on the image domain. Cremers et.al. have shown in [2] how to extend these models for scene segmentation. Recently we have shown how to use level set methods in conjunction with MRFs [4]. Global shape models are well suited e.g. for segmentation and tracking if the number of objects is known in advance and a good initial pose estimation is provided.

Semi global models consider shape characteristics in local neighbourhoods and go back to the ideas of G. Hinton on «product of experts» as well as of Roth and Black on «fields of experts» (see [6, 11] and citations therein). Mathematically these models are higher order GRFs of a certain type – additional auxiliary variables are used to express mixtures of local shape characteristics in usually overlapping neighbourhoods. Marginalisation over these auxiliary variables results in GRFs of higher order. The work of Kohli, Torr et.al. [10, 8] demonstrates how to introduce such higher order Gibbs potentials directly and to use them for segmentation in hierarchical Conditional Random Fields (CRF). However, it is not clear how to learn the graphical structure for such models.

Contributions

We will show that Gibbs Random Fields of the second order have already sufficient expressive power to model complex shapes as coherent spatial compositions of simpler parts. Obviously, these models have to have a significantly more complex graphical structure than just simple lattices. Moreover, the graphical structure itself be-
comes a parameter which has to be learnt together with the Gibbs potentials for each considered shape class.

From the application point of view these models have advantages especially in the context of scenes with an unknown number of similar objects (i.e. all objects are instances of a single shape class). Moreover, such models can be easily combined for scenes with instances of different shape classes.

The structure of the paper is as follows. In section 2 we introduce the GRF model for composite shapes and discuss the inference and learning tasks. The latter means to learn the Gibbs potentials and the graphical structure itself. The section 3 gives experiments exploring the expressive power of the model — first we separately show its ability to express spatial relations between segments and its ability to model simple shapes. Then we demonstrate its capability to model composite shapes including structure learning. Finally, we show how to combine such models for the discrimination of shape classes.

2. The shape model

Probability distribution

We begin with the description of the prior part of our shape model. Let \( D \subset \mathbb{Z}^2 \) be a finite set of nodes \( t \in D \), where each node corresponds to an image pixel. Let \( A \subset \mathbb{Z}^2 \) be a set of vectors used to define a neighbourhood structure on the set of nodes, i.e. a graph: two nodes \( t, t' \) are connected by an edge if \( t' - t = a \in A \). To avoid double edges we require \( -A \cap A = 0 \) (we use unary potentials as well). The resulting graph is obviously translational invariant and the elements of \( a \in A \) define subsets \( E_a \subset E \) of equivalent edges, where \( e = (t, t') \in E_a \) if \( t' - t = a \). A simple example is shown in Fig. 1.

Given a class of composite shapes, we denote the set of its parts enlarged by an extra element for the background by \( K \). A shape-part labelling \( y : D \rightarrow K \) is a mapping, that assigns either a shape-part label or the background label \( k \in K \) to each node \( t \in D \). A function \( u_a : K \times K \rightarrow \mathbb{R} \) is defined for each difference vector \( a \in A \). Its values \( u_a(k, k') \) are called Gibbs potentials. A corresponding probability distribution is defined over the set of shape-part labellings as follows:

\[
p(y) = \frac{1}{Z(u)} \exp \sum_{a \in A} \sum_{t \in E_a} u_a(y_t, y_{t'}),
\]

where \( Z \) denotes the partition sum (we omit the unary terms for better readability). This p.d. is homogeneously parametrised — all edges in an equivalence class \( E_a \) have the same potentials.

\[
\text{Fig. 1. Left: example of a translational invariant graphical structure.}
\]

Equivalence classes of edges \( E_a \) are coloured by different colours. The set \( A \) is represented by bold edges outgoing from the central node. Right: Gibbs potentials for an edge from \( E_a \).

Note that the parameters \( u_a \) of this model are unique up to additive constants for a given p.d. under fairly general assumptions — the only possible equivalent transformations (aka re-parametrisations) consist in adding a constant \( \tilde{u}_a() = u_a() + \text{const} \). This will be shown in appendix A.

Therefore, we assume from here that the Gibbs potentials for each \( a \in A \) are normalised to sum to zero: \( \sum_{k, k'} u_a(k, k') = 0 \).

It is important to notice that a homogeneously parametrised GRF on a finite domain \( D \subset \mathbb{Z}^2 \) is not necessarily homogeneous. A p.d. \( p(y) \) for labellings \( y : D \rightarrow K \) is called homogeneous if its marginals for congruent subsets coincide. This inhomogeneity, if present, usually reveals at the domain boundary. It is easy to verify that the converse is true at least for chains: a homogeneous Markov model on a finite chain admits a homogeneous parametrisation.

The appearance model is assumed to be a «simple» conditional independent model. The probability to observe an image \( x : D \rightarrow C \) (\( C \) is some colour space) given a shape-part labelling \( y \) is

\[
p(x | y) = \prod_{t \in D} p(x_t | y_t).
\]
In the light of the current popularity of CRFs it might well be asked, why we decided to favour a GRF here. Both variants are identical with respect to inference. Differences occur for learning. We can imagine that shape-part labellings can be used as latent variable layers for complex object segmentation models. Recently, empirical risk minimisation learning has been suggested for structured SVM models with latent variables [13]. This shows that the learning of graphical models with latent variables is possible for both variants − GRFs and CRFs. However, since we want to study the expressive power of the model in its pure form, we need a prior p.d. and moreover, we want to be able to learn such models fully unsupervised, which is possible for GRFs but not for CRFs.

**The inference task**

Informally, the inference task can be understood as follows. Given an observation (i.e. an image), it is necessary to assign values to all hidden variables. We pose the segmentation task as a Bayesian decision task. Let \( y' \) be the true (but unknown) segmentation and \( C(y, y') \) be a loss function, that assigns a penalty for each possible decision \( y \). The task of Bayesian decision is to minimise the expected loss

\[
R(y; x) = \sum_{y'} p(y' | x) C(y, y') \rightarrow \min_y.
\]

We use the number of misclassified pixels

\[
C(y, y') = \sum_t 1\{y_t \neq y'_t\}
\]

as the loss function. It leads to the max-marginal decision

\[
y_t^* = \max_k p(y_t = k | x) \quad \forall t \in D.
\]

Hence, it is necessary to calculate the marginal posterior probabilities for each node \( t \in D \) and label \( k \in K \). Currently this task is infeasible for GRFs. Several approximation techniques based e.g. on belief propagation or variational methods have been suggested for this task (see e.g. [12] for an overview). Unfortunately none of them guarantees convergence to the exact values of the sought-after marginal probabilities. To our knowledge, the only scheme which does it is sampling, which is however known to be slow [3].

**Estimation of Gibbs potentials**

The learning task comprises to estimate the unknown model parameters given a learning sample. We assume that the latter is a random realisation of i.i.d. random variables, so that the Maximum Likelihood estimator is applicable.

The following situations are distinguished depending on the format of the learning data. If the elements of the sample have the format \((x, y)\) then the learning is called supervised. If, instead, they consist of images only then the learning is called fully unsupervised. To cope with variants in-between as well, i.e. partial labellings \( y'_v \), we consider the elements of the training sample to be events of the type \( B = (x, y_v) = \{(y, x) | y'_v = y_v\} \).

We start with the learning of unknown potentials \( u \). For simplicity we consider the case when only one event \( B \) is given as the training sample. According to the Maximum Likelihood principle, the task is

\[
p(B; u) = \sum_{y \in B} p(y) p(x | y) \rightarrow \max_u.
\]

Taking the logarithm and substituting the model (1), (2) gives

\[
L(u) = \log \sum_{y \in B} \exp(\sum_{a \in A} \sum_{a' \in E_a} u_a(y_t, y_{t'}) p(x | y) - \log(Z(u)) \rightarrow \max_u.
\]

It is easy to prove, that the derivative with respect to the potentials is a difference of expectations of some random variable \( a_t k k' y_t y_{t'} \) with respect to the posterior and prior p.d.

\[
\frac{\partial L}{\partial u_a(k, k')} = \mathbb{E}_{p(y | B; u)}[n_a(k, k'; y)] - \mathbb{E}_{p(y; u)}[n_a(k, k'; y)].
\]

The random variables \( n_a(k, k'; y) \) are defined by

\[
n_a(k, k'; y) = \sum_{t \in E_a} 1\{y_t = k, y_{t'} = k'\}
\]

and represent co-occurrences for label pairs \((k, k')\) along the edges in \( E_a \) for a labelling \( y \). Combining these random variables into a random vector \( \Phi \), the gradient of the log-likelihood can be written as

\[
\nabla L(u) = \mathbb{E}_{p(y | B; u)}[\Phi] - \mathbb{E}_{p(y; u)}[\Phi].
\]
The exact calculation of the expectations in (8) is not feasible. Therefore, we suggest to use a stochastic gradient ascent to maximise (7). The learning algorithm is an iteration of the following steps:

1. Sample \( \tilde{y} \) and \( y \) according to the current a-posteriori probability \( p(y \mid \mathcal{B}; u) \) and a-priori probability \( p(y; u) \) respectively.

2. Compute \( n_a(k,k'; \tilde{y}) \) and \( n_a(k,k'; y) \) by (9) for each \( a \in A, k, k' \in K \).

3. Replace the expectations in (8) by their realisations and calculate new potentials \( u \).

For the sake of completeness we would like to mention that the learning of the appearance models \( p(c \mid k) \) can be done in a very similar manner. It is even simpler from the computational point of view because the normalising constant \( Z \) does not depend on these probabilities. Therefore it is not necessary to sample labellings according to the a-priori probability distribution \( p(y) \). Only a-posteriori sampled labellings are needed to perform the corresponding stochastic gradient step.

**Estimation of the interaction structure**

A very important question not discussed so far is the optimal choice of the neighbourhood structure \( A \). Unfortunately, no well founded answer to this question is known at present. One option is to use an abundant set of interaction edges, e.g. to use an exhaustive search over all possible sets \( A \) of given size \( |A| = m \) according to the Maximum Likelihood principle \( L(u_A, A) \rightarrow \max_{u_A, A} \). This task is however not feasible – an exhaustive search over all possible sets \( A \) would be computationally prohibitive, and, moreover, the likelihood can be calculated only approximatively. Therefore we rely on a greedy approximation which we will consider in two variants – one of them successively includes new elements into the neighbourhood structure starting from \( A = \{0\} \) and the other successively removes elements from this structure starting from \( A = \{a \in \mathbb{Z}^2 \mid |a_1| \leq d_1, |a_2| \leq d_2\} \).

For the first variant we use a greedy search for the interaction edges suggested by Zalesny and Gimel’farb in the context of texture modelling [14, 5]. Starting from the set \( A = \{0\} \), i.e. a model with unary potentials, new edges are iteratively chosen and included into \( A \) as follows. First, the optimal set of potentials \( u_A^* \in \mathcal{U}_A \) is determined for the current set \( A \) as described in the previous subsection. Here \( \mathcal{U}_A \) denotes the subspace of potentials on the edges in \( A \) (we may assume that the Gibbs potentials are zero on all other edges). If a bigger neighbourhood \( A' \) is considered, then clearly, the gradient of the (log) likelihood with respect to \( u_{A'} \) in the point \( u_A^* \) will be orthogonal to the subspace \( \mathcal{U}_A \). The proposal is to include the vector \( a' \in A' \setminus A \) with the largest gradient component

\[
\begin{align*}
\hat{a}' &= \arg \max_{a' \in A' \setminus A} \sum_{k,k'} [n_a(k,k'; \mathcal{B}, u) - n_a(k,k'; u)]^2. \\
&= \arg \max_{a' \in A' \setminus A} \sum_{k,k'} [n_a(k,k'; \mathcal{B}, u) - n_a(k,k'; u)]^2. \\
&\end{align*}
\]

Optionally the Kullback–Leibler divergence can be used instead of the Euclidean distance.

The second variant of structure estimation proceeds in opposite order. Starting with the neighbourhood structure \( A = \{a \in \mathbb{Z}^2 \mid |a_1| \leq d_1, |a_2| \leq d_2\} \), elements of \( A \) are successively removed. The aim is to remove in each step the element with the smallest impact on the maximal likelihood

\[
\max_{u_A} L(u_A) - \max_{u_{A,\hat{a}}} L(u_{A,\hat{a}}) \rightarrow \min, \quad (12)
\]

It is impossible to estimate this expression in the point \( u_A^* = \arg \max_{u_A} L(u_A) \) using the gradient of the likelihood (like in the first variant) because of \( \nabla L(u_A^*) = 0 \). It is nevertheless possible to estimate this expression based on \( u_A^* \). For the sake of simplicity we show this for the situation of supere-
vised learning. The likelihood maximisation with respect to the Gibbs potentials reads
\[
\max_{u_A} \{ <\psi_A, u_A> - \log \sum_y \exp <\phi_A(y), u_A> \} \tag{13}
\]
for this case. Here we have used the following notations. The set of all Gibbs potentials \( u_a(.,.) \), \( a \in A \) is considered as a vector \( u_A \). A realisation of the random vector \( \Phi_A \) (see (10)) is denoted by \( \phi_A(y) \). Finally, \( \psi_A \) denotes the corresponding vector of statistics resulting from the training sample. Designating \( \log Z(u_A) \) by \( H(u_A) \), the expression in (13) is nothing but the Fenchel conjugate \( H^*(\psi_A) \). It is known that for exponential families the latter can be written as
\[
\psi_A^* = \inf\{ \log p(y) | \mathbb{E}_p[\Phi_A] = \psi_A \}, \quad p \in \mathbb{E}
\tag{14}
\]
(see e.g. [12, 1]), where we denoted the expectation w.r.t. a probability distribution \( p \) by \( \mathbb{E}_p \) and the set of all probability distributions on labellings \( y \) by \( \mathbb{P} \). This means to find the p.d. with maximal entropy among all distributions having expectation \( \psi_A \) of the random vector \( \Phi_A \).

Removing an element \( a \) from the neighbourhood structure \( A \) can be equivalently expressed by the linear constraints \( u_a = 0 \). Considering the task (13) with these additional constraints, it can be shown by the use of Fenchel duality (see e.g. [1]) that the corresponding conjugate function \( \tilde{H}^*(\psi_A) \) can be written as
\[
\tilde{H}^*(\psi_A) = \inf_{z_a} \{ \sum_y p(y) \log p(y) | \mathbb{E}_p[\Phi_A] = \psi_A + z_a \}, \quad p \in \mathbb{E}
\tag{15}
\]
where \( z_a \) is an arbitrary vector of the subspace \( \mathcal{U}_a \). Therefore, the difference in (12) is equal to
\[
H^*(\psi_A) - \tilde{H}^*(\psi_A) = H^*(\psi_A) - H^*(\psi_A + z_a^*)
\]
and can be estimated by the gradient of \( H^* \) in \( \psi_A \). The latter gradient is nothing but the vector of Gibbs potentials \( u_A^* \).

The convex, lower semi-continuous function \( H^*(\psi_A) \) is not differentiable in general. Therefore its sub-differential may consist of more than one subgradient \( u_A \). This corresponds to the non-uniqueness of the Gibbs potentials. We have however shown that the Gibbs potentials are unique up to additive constants for the model class considered in this paper (see Appendix A).

Summarising, the difference in (12) can be estimated by \( \|u_a\| \), that leads to the following greedy removal strategy for elements of the neighbourhood structure \( A \). Given a current neighbourhood structure \( A \), estimate the optimal Gibbs potentials \( u_A^* \) and remove the element \( a \in A \) with the smallest value of \( \|u_a\| \).

3. Experiments

Modelling spatial relations between segments

The first experiment investigates the ability of the model (1), (2) to reflect spatial relations between segments, i.e. scene parts, which are too large to capture their shape by a neighbourhood structure of reasonable size. We used the three images shown in the first row of Fig. 2 as training examples. Each scene should be segmented into three segments: \( K = \{ \text{sky}, \text{trees}, \text{grass} \} \). The appearance models \( p(c|k) \) for the segments were assumed as mixtures of multivariate Gaussians (four per segment). A model with «full» neighbourhood structure — all vectors \( \{ a \in \mathbb{Z}^2 | a_i \leq d, |a_2| \leq d \} \) with \( d = 20 \) was used in this experiment. A «simple» but anisotropic Potts model on the 8-neighbourhood was chosen as a baseline for comparison.

Semi-supervised learning was applied by fixing the segment labels in the rectangular areas shown by rectangles during learning. Both the a-priori models (the potentials and the direction specific Potts parameters for the baseline model) and the appearance models (mixture weights, mean values and covariance matrices) were learned.

The difference of the models can be clearly seen by observing labellings generated a-priori by the learned models, i.e. without input images. Some of them are shown in the second and third row for the model with complex neighbourhood structure and the baseline model respectively. It
can be seen, that the spatial relations between segments (like e.g. «above», «below» etc.) were correctly captured by the complex model, whereas it is clearly not the case for the Potts model.

The consequences can be clearly seen from the following experiment. We fixed the prior models obtained in the previous experiment (semi-supervised learning) for both variants (the complex prior and the Potts prior) and learned the parameters of the Gaussian mixtures completely unsupervised. Fig. 3 shows labellings (i.e. segmentations) sampled at the end of the learning process by the corresponding a-posteriori probability distributions (obtained with the learned appearances) for the complex a-priori model and the Potts a-priori model in the first and the second row respectively. The advantages of the complex model are clearly seen. These results can be explained as follows. There are twelve Gaussians in total to interpret the given images. For the learning process it is «hard to decide» which of the Gaussians belongs to which segment. Using the compactness assumption only is obviously not enough to separate segments from each other. If the complex model is used instead, the learning process starts to generate labellings according to the a-priori probability distribution, i.e. labellings which reflect the correct spatial relations between the segments. This forces the unsupervised learning of the appearance models into the right direction.

**Modelling simple shapes**

This group of experiments demonstrates the ability of the model to represent simple shapes as well as to perform shape driven segmentation. This experiment is prototypical e.g. for a class of image recognition tasks in biomedical research. Fig. 4 (upper left) shows a microscope image of liver cells with stained DNA. Thus, only the cell nuclei are visible. The task is to segment the image into two segments — «cells» (which have nearly circular shape) and «background» (the rest...
including artefacts). Hence, two labels are used. The «full» neighbourhood structure with \( d = 12 \) was used (it approximately corresponds to the mean cell diameter). Again, we used a baseline model for comparison – a GRF with 4-neighbourhood and free potentials. The appearances for grey-values were assumed to be Gaussian mixtures (two per segment) in both models.

\[ u(k, k') = \begin{cases} \alpha & \text{if } k = k', \\ -\alpha & \text{else.} \end{cases} \quad \text{(16)} \]

The Gibbs potentials on the short vectors are supermodular and express the correlation of the labels on the edges of this type

\[ u_2(k, k') = \begin{cases} \beta & \text{if } k = k' = 0, \\ -\beta & \text{if } k = k' = 1, \\ 0 & \text{else,} \end{cases} \quad \text{(17)} \]

is used to influence the density of the blobs. A labelling (fragment) sampled by this model \((\alpha = 0.35, \beta = 0.5)\) is shown in Fig. 5. Both heuristic approaches for structure estimation discussed in the previous section were applied for the supervised version, i.e. using a labelling generated by the known model as a learning sample.
The first approach−iterative growth of the structure−was run 40 times. The estimated structures resulting from these runs are shown in Fig. 5 as a grey-coded histogram. As a stochastic gradient ascend is used for the learning of the potentials, each run may result in a different structure. The histogram shows however, that the structure estimation is essentially correct. All trials of the second approach−iterative shrinking of the neighbourhood structure−resulted much to our surprise in one and the same estimated structure−the correct one.

We conclude from these experiments that the neighbourhood structure of a shape model is identifiable (at least in principle) from labellings generated by the model.

Modelling composite shapes

The previous experiments have shown that second order GRFs can model both, spatial relations between segments and simple shapes. Now we are going to demonstrate the capability of the model to capture both properties simultaneously. This opens the possibility to represent complex shapes as spatial compositions of simpler parts.

To demonstrate this, we use an artificial example shown in Fig. 6 (upper left). It was produced manually and corrupted by Gaussian noise. Accordingly, the model was defined as follows. The label set $K$ consists of seven labels, each one corresponding to a part of the modelled shape (as well as one for the background). The appearance models $p(c|k)$ for the labels are Gaussians with known parameters. In this experiment we applied the growth variant for the estimation of the interaction structure as described in section 2.

Fig. 6 (upper row, center) shows a labelling generated by the learned prior model. It is clearly seen that both, spatial relations between object parts and part shapes are captured correctly.

The bottom row of Fig. 6 displays labellings generated during the process of structure learning at time moments, when the interaction structure learned so far was not yet capable to capture all needed properties. As it can be seen, the model was able to learn spatial relations between the segments more or less correctly even for a small numbers of edges (5 edges−bottom left). More relations are learned as the number of edges grows (bottom middle and right). Finally, 20 difference vectors were necessary to capture all relations (out of 1200 possible for the maximal range of $d = 24$).

Fig. 6 (upper right) shows the estimated neighbourhood structure. The endpoints of all edges from central pixel are marked by colours (the image is magnified for better visibility). A certain structure can be seen in this image. The 8-neighbourhood edges (black) reflect compactness and adjacency relations of the object parts. The learned potentials on these edges represent strong label co-occurrences. Most of the other vectors are
responsible for the shapes of the parts. The potentials on the red edges express characteristic breadths, and the potentials on the green edges – characteristic lengths of the parts. The potentials on these edges mainly represent anti-correlations, forcing label values to change along certain directions. The blue pixels in the figure reflect relative positions of object parts.

**Composite shape recognition**

The final experiment demonstrates possibilities to combine composite shape models. The aim is to obtain a joint model which can be used for detection, segmentation and classification of objects in scenes populated by instances of different shape classes like e.g. the example in Fig. 7. We conclude from the previous experiments, that the appearance model can be re-learned in a fully unsupervised way if the prior shape model is discriminative. Hence, the most important question is, how to combine the prior models. We propose a method for this that is based on the following observation. It is not necessary to have an example image (or an example segmentation) in order to learn the model if the aposteriori statistics

\[
\Phi_a(k,k') = \mathbb{E}_{p(y|b,a)}[\phi_a(k,k')]
\]  

(18)

for all difference vectors \(a \in A\) and label pairs \((k,k')\) are known – the gradient of the likelihood (equation (8)) reads then

\[
\frac{\partial L}{\partial u_a(k,k')} = \Phi_a(k,k') - \mathbb{E}_{p(y,a)}[n_a(k,k';y)] .
\]  

(19)

Let us consider this in a bit more detail for a simple example – just two shapes like in Fig. 7. Let us assume that the both models are learned, i.e. both the potentials and statistics are known for both models and for all difference vectors \(a\). Obviously, it is not easy to combine the potentials of both shape models in order to obtain new ones for a model that generates such collages. It is however very easy to estimate the needed aposteriori statistics for the joint model given the aposteriori statistics for both shape models. Summarizing, the scheme to obtain the parameters of the joint model consists of two stages:

1. compute the aposteriori statistics for the joint model and

2. learn the model according to (19) so that it reproduces this statistics.

As the second stage is standard, we consider the first one in more detail. Let us denote the label sets corresponding to the shape parts by \(K_1\) and \(K_2\) for the first and for the second shape type respectively. Let \(b_1\) and \(b_2\) be the background labels in the corresponding shape models and \(b\) be the background label in the joint one. Consequently, the label set of the latter is \(K_1 \cup K_2 \cup b\) (see the middle part of Fig. 8).

First of all we enlarge the label sets of each shape model by labels that are not present in this model but present in the joint one. Thereby the statistics for the new introduced labels (for all difference vectors \(a\)) are set to zero (see Fig. 8, left and right). Informally said, these extended aposteriori statistics correspond to the situations that the joint model is learned on examples, in which only labels of one particular shape are present. The aposteriori statistics for the joint model is then obtained as a weighted mixture of the two extended ones and an additional uniformly distributed component. The latter is added in order to avoid zero probabilities (which would lead to obvious techni-
cal problems for the Gibbs Sampler). Summarising, the a posteriori statistics of label pairs for a difference vector $a$ of the joint model is:

$$
\Phi_a(k,k') = \begin{cases} 
  w_1 \cdot \Phi_a(k,k') + w_0 & \text{if } k \in K^1 \text{ and } k' \in K^1, \\
  w_1 \cdot \Phi_a(k,k') + w_0 & \text{if } k \in K^2 \text{ and } k' \in K^2, \\
  w_1 \cdot \Phi_a(k,k') + w_0 & \text{if } k = b \text{ and } k' = b, \\
  w_1 \cdot \Phi_a(b^1,b^1) + w_2 \cdot \Phi_a(b^2,b^2) + w_0 & \text{if } k = b \text{ and } k' = b \\
  w_0 & \text{otherwise.}
\end{cases} 
$$

(20)

with some weights $w_0 \ll w_1 \approx w_2$, where the indices 1 and 2 correspond to the particular shape model. Given these statistics the joint model is learned according to (19).

For the experiment in Fig. 7 two composite shape models were learned separately. The test image in Fig. 7 (left) is a collage of both shape types. Note that the appearance of all shape parts is identical, so they are not distinguishable without the prior shape model. Fig. 7 (right) shows the final segmentation. It is seen that all objects were correctly segmented and recognised – although both composite shape classes share some similarly shaped parts – they were not confused.

4. Conclusions

The notation of shape is often understood as an object property of global nature. We followed a different direction by modelling shapes in a distributed way. We have demonstrated that the expressive power of second order GRFs allows to model spatial relations of segments, simple shapes and moreover, both aspects simultaneously i.e. composite shapes which are understood as coherent spatial compositions of simpler shape parts.

We have shown that complex shapes can be recognized even in the situation, when their parts are not distinguishable by appearance. However, in our learning experiments we used training images, where they are distinguishable. Thus, an important question is, whether it is possible to perform unsupervised decomposition of complex shapes into simpler parts during the learning phase, i.e. to learn shape models from images, where the desired spatial relations between shape parts are not explicitly present. Another important issue is the learning of the interaction structure. It would be very useful to have a well grounded approach for this.

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can be equivalently written as

with components \( \omega \) are to \( - \sum \omega \). Consequently, (30)

\( \omega \), then, clearly, considering equation (27) for each of \( \omega \) privacy for shape part labellings

\( \mathcal{A} \). We call two parametrisations \( \omega \) equivalent, if the corresponding probability distributions are identical. It follows that the difference \( \omega = \omega - \omega \) of equivalent potentials fulfils

\[
V(y) = \sum_{t} v_{t}(k) n_{t}(k; y) + \left[ -\sum_{t \in T} \sum_{k} v_{t}(k, k') n_{t}(k, k'; y) \right] = \text{const.}
\]

We will conclude that all functions \( v_{t} \) are constant under fairly general conditions. We perform the proof in two steps. First we show that the pairwise functions \( v_{t}, \quad a \neq 0 \) are modular and can be written as a sum of unary functions. In a second step we will conclude the claimed statement under fairly general conditions for the graph \( (D, E) \).

Let us consider an arbitrary non-zero vector \( a \epsilon A \) of the neighbourhood structure and an arbitrary edge \( (tt') \in E_{a} \). Let \( k_{1}, k_{2} \) be two arbitrary labels in the node \( t \) and \( k_{1}', k_{2}' \) be two arbitrary labels in the node \( t' \). Let \( y_{11}, y_{12}, y_{21}, y_{22} \) be four labellings with respective values \( (k_{1}, k_{1}'), (k_{1}, k_{2}'), (k_{2}, k_{1}'), (k_{2}, k_{2}') \) on the nodes \( t, t' \) such that they coincide on all other vertices. We consider the equation

\[
V(y_{11}) + V(y_{12}) - V(y_{21}) - V(y_{22}) = 0.
\]

It is easy to see that this equation reduces to

\[
v_{t}(k_{1}, k_{1}') + v_{t}(k_{2}, k_{2}') - v_{t}(k_{1}, k_{1}') - v_{t}(k_{2}, k_{2}') = 0.
\]

This holds for arbitrary four-tuples of labels and it follows that the function \( v_{t} \) is modular and can be written as a sum of two unary functions

\[
v_{t}(k, k') = \tilde{v}_{t}(k) + \hat{v}_{t}(k').
\]

These arguments can be applied for every element \( a \in A' \). Consequently, \( V(y) \) can be written as

\[
V(y) = \sum_{k} v_{a}(k) n_{a}(k; y) + \frac{1}{2} \sum_{a \in A} \sum_{k} [v_{a}(k) n_{a}(k; y) + \hat{v}_{a}(k) n_{a}(k; y)],
\]

where we have omitted the tildes. Note that \( n_{a}(k; y) = \sum_{k} n_{a}(k, k'; y) \) denotes the number of vertices with an outgoing edge of type \( a \) for which the labelling \( y \) has the value \( k \). Therefore in general \( n_{a}(k; y) \neq n_{a}(k; y') \).

Let us consider an arbitrary vertex \( t \) and two labellings \( y, \bar{y} \) which coincide on all vertices but \( t \). It follows from \( V(y) = V(\bar{y}) = 0 \) that

\[
v_{t}(k) + \sum_{a \in D} v_{a}(k) + \sum_{a \in D} \hat{v}_{a}(k) = \text{const.}
\]

We assign a vector \( z(t) \) with dimension \( 2|A| - 1 \) to every vertex \( t \in D \) with components

\[
z_{t}(a) = \begin{cases} 1 & \text{if } a \in D, \\ 0 & \text{else.} \end{cases}
\]

If the domain \( D \) contains a subset of nodes \( t \) such that their vectors \( z(t) \) span the whole vector space of dimension \( 2|A| - 1 \), then, clearly, considering equation (27) for each of them, we obtain

\[
v_{t}(k) = \text{const.}
\]

\[
v_{a}(k) = \text{const.}
\]

\[
\hat{v}_{a}(k) = \text{const.}
\]

for all \( a \in A \).

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