Convexity Shape Prior for Binary Segmentation
Lena Gorelick, Olga Veksler, Yuri Boykov, and Claudia Nieuwenhuis

Abstract—Convexity is a known important cue in human vision. We propose shape convexity as a new high-order regularization constraint for binary image segmentation. In the context of discrete optimization, object convexity is represented as a sum of 3-clique potentials penalizing any 1-0-1 configuration on all straight lines. We show that these non-submodular potentials can be efficiently optimized using an iterative trust region approach. At each iteration the energy is linearly approximated and globally optimized within a small trust region around the current solution. While the quadratic number of all 3-cliques is prohibitively high, we design a dynamic programming technique for evaluating and approximating these cliques in linear time. We also derive a second order approximation model that is more accurate but computationally intensive.

We discuss limitations of our local optimization and propose gradual non-submodularization scheme that alleviates some limitations. Our experiments demonstrate general usefulness of the proposed convexity shape prior on synthetic and real image segmentation examples. Unlike standard second-order length regularization, our convexity prior does not have shrinking bias, and is robust to changes in scale and parameter selection.

Index Terms—Segmentation, convexity shape prior, high-order functionals, trust region, graph cuts.

I. INTRODUCTION

Length-based regularization is commonly used for ill-posed segmentation problems, in part because efficient global optimization algorithms are well-known for both discrete and continuous formulations, e.g. [3], [23]. Nevertheless, the shrinking bias and the sensitivity to the weight of the length term in the energy are widely recognized as limitations of this form of regularization. These problems motivate active research on optimization of higher-order regularization energies, e.g. curvature [24], [6], [22], [20] and cooperative prior [12], [13], which can alleviate the shrinking bias and other issues.

We propose a new higher-order regularization model: convexity shape constraint, see Fig. 1. Convexity was identified as an important cue in human vision [18], [19]. Many natural images have convex or nearly convex objects. Convex objects are also common in medical images. Yet, to the best of our knowledge, we are the first to introduce a convexity shape prior into discrete segmentation energy.

We develop an energy-based formulation for convexity prior in discrete optimization framework and propose an efficient optimization algorithm for the corresponding non-submodular high-order energy term. For a given segment $S \subset \Omega$, the overall segmentation energy $E(S)$ can combine our convexity prior $E_{convexity}(S)$ with user-defined hard-constraints, linear appearance models [4], boundary length [3], color separation [26], or any others standard submodular terms $E_{sub}(S)$

$$E(S) = E_{convexity}(S) + E_{sub}(S).$$

Convexity of segment $S$ is expressed as a penalty for all ordered triplet configurations 1-0-1 along any straight line, see Fig. 2. Similar straight 3-cliques also appear in curvature modeling [20], but they also need 0-1-0 configurations to penalize negative curvature. Moreover, they use only local triplets to evaluate curvature. In contrast, convexity is not a local property of the segment boundary. Therefore, we have to penalize 1-0-1 configurations on straight intervals of any length. Consequently, our convexity energy model has a much larger number of cliques. We propose an efficient dynamic programming technique to evaluate and approximate these cliques in the context of trust region optimization [10].

Related Work: Many related shape priors were introduced in the past. Common length-based regularizer [3] penalizes segment perimeter favoring smooth solutions that are closer to circles and, therefore, more convex. However, as shown in our experiments, this prior needs to be carefully balanced with the appearance term as it has a strong shrinking bias. Connectivity regularizer [28], [21] does not have shrinking bias but might suffer from connecting thin structure artifacts.

Another related regularizer is the star shape prior [27], [11], which imposes convexity constraints only along the lines passing through a reference point given by the user: these lines are allowed to enter and exit the object only once. In contrast to our convexity prior, the star shape prior allows for non-convex objects, e.g. a star.

There are also part-based shape priors [29], [17], [7]. A shape is partitioned into several parts and each part imposes certain constraints on the direction of the boundary with the background. This approach can model some simple convex shapes, e.g. a rectangle, but it can not represent a general convexity prior.

The most related work to ours is [25], which models the object as an $n$-sided convex polygon. It is a part-based approach that uses one foreground and $n$ background labels. For an accurate segmentation of an arbitrary convex object, e.g. a circle, a finer discretization (i.e. more background parts) is required, significantly increasing runtime. The larger the object, the worse is the problem. In contrast, we can obtain an arbitrary convex object for any choice of orientation discretization. Moreover, [25] relies on continuous optimization and is not efficient without GPU. Additional related work on optimization is discussed in Sec. IV-C.

Contributions: We introduce a new discrete convexity shape regularizer. When enforced as a hard constraint, it is a parameter-free convexity shape prior. In practice, we enforce convexity as a soft constraint using a finite penalty.

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Manuscript received June XX, 2015; revised Month XX, 2015.
However, our experiments show that there is almost no variation in segmentation results for different values of the convexity weight $\omega$, once the value is high enough. Our method is also robust to changes in scale, see Sec. IV.

We develop an optimization algorithm based on trust region framework and show how to use dynamic programming to significantly improve efficiency. In the context of trust region, we derive and compare both linear and quadratic approximations of our energy. While quadratic approximation is more accurate, it is computationally more expensive. Nonetheless, it could potentially be useful for approximating other higher-order energies.

We discuss alternative convexity models such as central clique model and local convexity model and alternative optimization schemes such as gradual non-submodularization. We also experimentally validate the advantage of our convexity vs. the length regularizer for segmentation.

Finally, we collected a new dataset of natural images with ground truth convex segmentations, available for download.

The paper is organized as follows. Sec. II formulates convexity energy and Sec. II-A explains its efficient evaluation. Sec. III introduces trust region optimization framework. We derive linear approximation of our energy in Sec. III-A and show how to compute it efficiently using dynamic programming in Sec. III-B. In Sec. III-C we discuss a more accurate quadratic approximation of our energy. Sec. IV-A introduces trust region optimization framework and show how to use dynamic programming to compute it efficiently. Sec. IV-B discusses alternative optimization schemes such as convexity models and alternative orientations, see Fig. 2 (left). Let $p \in \Omega$ be a pixel, and let $l^i_p$ be a discrete line passing through $p$ in orientation $d_i$. That is,

$$l^i_p = \{ p t \}$$

We define $L_i = \{ l^i_p \in \Omega | p \in \Omega \}$ as the set of all discrete lines $l^i_p$ of given orientation $d_i$. Fig. 2 (middle) illustrates set $L_i$ for one particular orientation $d_i$ and highlights one $l^i_p$. To avoid double indexing throughout the paper we use $x_i$ instead of $x_{p_t}$ to denote the binary variable for pixel $p_t$ on a line $l^i_p$.

One way to represent discrete convexity constraint is based on potential $\phi : \{0, 1\}^3 \rightarrow \mathbb{R}$ defined for all triplets of ordered pixels $(p_s, p_t, p_v)$, $s < t < v$ along any discrete line $l \in \bigcup L_i$

$$\phi(x_s, x_t, x_v) = \begin{cases} \infty & \text{if } (x_s, x_t, x_v) = (1, 0, 1) \\ 0 & \text{otherwise.} \end{cases}$$

In practice we use some finite penalty $\omega$ redefining potential $\phi$ algebraically as

$$\tilde{\phi}(x_s, x_t, x_v) = \omega \cdot x_s (1 - x_t) x_v.$$  \hfill (3)

The convexity energy $E_{\text{convexity}}(x)$ integrates this triple clique potential over all orientations, all lines and all triplets:

$$E_{\text{convexity}}(x) = \sum_{l \in \bigcup L_i} \sum_{(p_s, p_t, p_v) \in l} \tilde{\phi}(x_s, x_t, x_v).$$  \hfill (4)

As discussed below, 3rd-order energy (4) is hard to optimize for two reasons: it is non-submodular and it has a prohibitively large number of cliques.

It is easy to verify that this energy is non-submodular [8]. It is enough to show that there exist segments $S, T \subset \Omega$ s.t.

$$E(S) + E(T) < E(S \cap T) + E(S \cup T)$$  \hfill (5)

Note, pixel $p \in \Omega$ has unique index $t$ on each line $l$ passing through it.
Consider the example below. Since both $S$ and $T$ are convex, the left hand side in (5) is zero, while the right hand side is infinite since the union of $S$ and $T$ is not convex. Therefore, our energy cannot be optimized with standard methods for submodular functions.

At the first glance, is seems prohibitively expensive to even evaluate our energy on reasonably sized images. For example, for a 200 \times 300 image, with just 8 orientations, there are roughly 32 billion triple cliques. In Sec. II-A, III-B we show how to evaluate and approximate the $E_{\text{convexity}}$ in time linear w.r.t. image size using dynamic programming. Then, in Sec. III we show how to optimize our energy using trust region techniques [30], [10]. Alternative convexity models and optimization schemes are discussed in Sec. IV-B and IV-C.

### A. Energy Evaluation via Dynamic Programming

This section explains how to evaluate our convexity term $E_{\text{convexity}}(x)$ efficiently. We show how to compute the inner summation in (4) for one given line $l$. The idea is to use dynamic programming to efficiently count the number of triplets $(1,0,1)$ on a line violating convexity constraints.

Let $x_l$ denote a vector of binary indicator variables on line $l$. We rewrite

$$E_{\text{convexity}}(x_l) = \sum_{(p_s,p_t,p_v) \in l \atop s < t < v} \phi(x_s,x_t,x_v) = \omega \cdot \sum_{s < t < v} x_s \cdot (1 - x_t) \cdot x_v.$$  

Consider pixels $p_s,p_t,p_v \in l$. We say pixel $p_s$ precedes pixel $p_t$ on line $l$ if $s < t$. Similarly, pixel $p_v$ succeeds pixel $p_t$ if $v > t$. Let $C^-(t)$ be the number of pixels $p_s$ preceding pixel $p_t$ such that $x_s = 1$, and $C^+(t)$ be the number of pixels $p_v$ succeeding pixel $p_t$ such that $x_v = 1$:

$$C^-(t) = \sum_{s < t} x_s, \quad C^+(t) = \sum_{v > t} x_v.$$  

To count the number of all violating configuration $(1,0,1)$ for ordered triplets on line $l$ we first consider one fixed pixel $p_t \in l$ with zero label $x_t = 0$. Each preceding pixel with label one and each succeeding pixel with label one form configuration $(1,0,1)$. Thus, the total combinatorial number of ordered triplets $(p_s,p_t,p_v)$, $s < t < v$, with configuration $(1,0,1)$ is given by $C^+(t) \cdot C^-(t)$, see Fig. 3. Summing over all zero label pixels on line $l$ gives

$$E_{\text{convexity}}(x_l) = \omega \cdot \sum_t C^+(t) \cdot C^-(t) \cdot (1 - x_t).$$

Note that $C^-(t) = C^-(t-1) + x_{t-1}$ and $C^+(t) = C^+(t+1) + x_{t+1}$. Hence each of $C^+(t)$ and $C^-(t)$ can be computed for all pixels on a line in one pass using running sums. For a particular orientation $d_i$, each pixel appears in one line only. Therefore, the total number of operations needed to compute $E_{\text{convexity}}(x)$ is $O(mN)$, where $N = |\Omega|$ is the number of pixels in the image and $m$ is the number of distinct orientations.

![Fig. 3. Evaluation of $E_{\text{convexity}}$. The top row shows current configuration $x_l$ of pixels on line $l$. The second and the third rows show the number of pixels $p_s$ with $x_s = 1$ before and after each pixel $p_t$, that is, functions $C^-(t)$ and $C^+(t)$. The last row shows the number of violated constraints for each $p_t$ with $x_t = 0$, resulting in total of 12 violations on the line.](image-url)
Algorithm 1: Trust Region Convexity

1. \( x^0 \leftarrow x_{\text{init}}, \lambda_0 \leftarrow \lambda_{\text{init}}, \text{convergedFlag} \leftarrow 0 \)
2. Repeat Until convergedFlag
3. //Approximate \( E_{\text{convexity}}(x) \) around \( x^k \)
4. Compute \( E_{\text{approx}}^k(x) \) (see Sec. III-A)
5. \( E_k(x) = E_{\text{approx}}^k(x) + E_{\text{sub}}(x) \) // keep the submodular part
6. //Trust region sub-problem
7. \( x^* \leftarrow \arg\min_{x} L(x) \) (8)
8. Evaluate \( E_{\text{convexity}}(x^k), E_{\text{convexity}}(x^*) \) (see Sec. II-A)
9. Evaluate \( E_{\text{approx}}^k(x), E_{\text{approx}}^*(x^*) \) (see Sec. III-B)
10. \( R = E(x^k) - E(x^*) \) // actual reduction in energy
11. \( P = E^*(x^*) - E^*(x^k) \) // predicted reduction in energy
12. If \( P = 0 \) // meaning \( x^* = x^k \) and \( \lambda > \lambda_{\text{max}} \)
13. \( \lambda_k \leftarrow \lambda_{\text{max}} \)
14. //Trust region sub-problem:
15. \( x^* \leftarrow \arg\min_{x} L(x) \) (8)
16. Evaluate \( E_{\text{convexity}}(x^k), E_{\text{convexity}}(x^*) \) (see Sec. II-A)
17. Evaluate \( E_{\text{approx}}^k(x), E_{\text{approx}}^*(x^*) \) (see Sec. III-B)
18. \( R = E(x^k) - E(x^*) \) // actual reduction in energy
19. \( P = E^*(x^*) - E^*(x^k) \) // predicted reduction in energy
20. //Update current solution
21. \( x^{k+1} \leftarrow \begin{cases} x^* & \text{if } R > 0 \\ x^k & \text{otherwise} \end{cases} \)
22. convergedFlag \leftarrow \begin{cases} R \leq 0 & \text{if } \text{convergedFlag} \end{cases}
23. Else // meaning \( x^* \neq x^k \) and \( \lambda \leq \lambda_{\text{max}} \)
24. //Update current solution
25. \( x^{k+1} \leftarrow \begin{cases} x^* & \text{if } R > 0 \\ x^k & \text{otherwise} \end{cases} \)
26. //Adjust the trust region
27. \( \lambda_{k+1} \leftarrow \begin{cases} \lambda_k/\alpha & \text{if } R/P > \tau_2 \\ \lambda_k : \alpha & \text{otherwise} \end{cases} \)

we use \( \alpha = 10 \), \( \tau_2 = 0.25 \)

adaptively changed (line 27), based on the quality of the current approximation as motivated by empirical inverse proportionality relation between \( \lambda_k \) and \( d_k \) (see [10]). In each iteration of the trust region, either the energy decreases or the trust region size is reduced. When the trust region is so small that it does not contain a single discrete solution, namely \( x^* = x^k \) (Line 12), one more attempt is made using \( \lambda_{\text{max}} \), where \( \lambda_{\text{max}} = \sup \{ \lambda | x^* \neq x^k \} \) (see [10]). If there is no reduction in energy with smallest discrete step \( \lambda_{\text{max}} \) (Line 21), we are at a local minimum [5] and we stop (Line 22).

A. Linear Approximation of \( E_{\text{convexity}} \)

Below we derive linear approximation \( E_{\text{approx}}^k(x) \) for the energy term \( E_{\text{convexity}}(x) \) in (4) around current solution \( x^k \)

\[
E_{\text{approx}}^k(x) = \sum_{t \in \mathcal{L}} \sum_{x \in \mathcal{L}^k} \phi^k(x_s, x_t, x_v) \]

where \( \phi^k(x_s, x_t, x_v) \) is a linear approximation of the corresponding \( \phi(x_s, x_t, x_v) \) in (3) around \( x^k \), as explained below.

Property III-A.1. For any potential \( \phi(x) : \{0, 1\}^n \rightarrow \mathcal{R} \) of \( n \) binary variables \( x = (x_1, \ldots, x_n) \) and any subset \( A \subset \{0, 1\}^n \) of \( n + 1 \) distinct binary configurations of \( x \), there is a linear function \( L_A(x) = a_0 + a_1 x_1 + a_2 x_2 + \ldots + a_n x_n \) such that \( \phi(x) = L_A(x) \) for any \( x \in A \).

The proof of Prop. III-A.1 is by construction. To find linear approximation \( L_A(x) \) of any potential \( \phi(x) \), we need to compute \( n + 1 \) unknown coefficients of \( L_A(x) \). This can be done by solving a system of \( n + 1 \) equations \( \phi(x) = L_A(x) \) for \( n + 1 \) binary configurations in some chosen \( A \). Note that there are \( 2^n \) possible configurations of \( n \) binary variables. Therefore, for our triple clique potential in (3), set \( A \) contains four out of eight distinct configurations of labels \( (x_s, x_t, x_v) \). Since in our energy we have a large number of overlapping triple cliques, we apply property III-A to simultaneously approximate the energy on all triple cliques and sum up the resulting approximations.

In practice, for each triple clique, we use an approach that avoids solving systems of equations and implicitly selects a specific set \( A \). Note, any discrete potential \( \phi \) can be written as a combination of multilinear functions of variables \( (x_1, \ldots, x_n) \), see (3). In this case, it is easy to verify that Taylor expansion \( \phi_k \) of the potential \( \phi \) around configuration \( x^k \) is a linear function satisfying Prop. III-A.1. That is, \( \phi_k(x) \) agrees with \( \phi(x) \) on configuration \( x^k \) and other “neighboring” configurations obtained by flipping one of the variables in \( x^k = (x^k_1, \ldots, x^k_n) \).

In our case, omitting the constant terms, Taylor expansion of (3) around \( x^k \) yields:

\[
\phi_k(x_s, x_t, x_v) = (1 - x^k_t) \cdot x^k_v \cdot x_s \]

The components in (10) have an intuitive interpretation. Consider the first component \( (1 - x^k_t) \cdot x^k_v \cdot x_s \). Recall that pixels \( p_s, p_t, p_v \) are on a line and \( p_s \) is between \( p_t \) and \( p_v \). If the current configuration \( x^k \) is such that \( x^k_t = 0 \), and \( x^k_v = 1 \), then assigning label 1 to pixel \( p_s \) violates convexity, assuming \( p_s \) and \( p_v \) keep their labels unchanged from \( x^k \). The unary term \( (1 - x^k_t) \cdot x^k_v \cdot x_s \) penalizes this violation: assignment \( x_s = 1 \) carries a penalty, whereas \( x_s = 0 \) is not penalized. The other two components in (10) have similar intuitive interpretations.

Summing approximations of all triple potentials in (9) results in a linear function \( E_{\text{approx}} (x) \) that coincides with the original \( E_{\text{convexity}}(x) \) on current solution \( x^k \).

Approximation in (10) gives three unary terms for each triple clique. Consider pixel \( p_s \in l \). It can be either the leftmost, middle, or rightmost member of a clique on that line. We need to sum the terms from all triple cliques on line \( l \) involving pixel \( p_s \). First with \( p_s \) being on the left, then in the middle and finally on the right of the clique. All these terms contribute to the unary potential \( u^l_s(x_s) \) for a single pixel \( p_s \):
The full Taylor based unary term for pixel $p \in \Omega$ sums the above expression over all lines passing through $p$.

Fig. 5 illustrates the resulting unary terms arising from such approximation. They encourage any holes or concavities in the foreground segment to be filled in, and any protrusions/disconnected components to be erased. Efficient computation of (11) is discussed in Section III-B. There is a relation between the Taylor unary terms in (11) and parallel ICM algorithm, first noted in [16]. However, our trust region framework has many differences from parallel ICM [16]. See [9] for the experimental comparison.

B. Computation of $E_{\text{approx}}^k$ via Dynamic Programming

Naive computation of the summations in (11) is too costly. We now explain how to compute the unary terms in (11) efficiently. Similarly to Sec. II-A, the speedup is achieved with running sums on each line.

Let $s, t, v$ enumerate the pixels on line $l$. In (11), the first sum counts the number of pixel pairs $(p_s, p_v)$ such that $s < t < v$ and $x_s^k = 0$, $x_v^k = 1$. The second sum counts the number of pixels pairs $(p_s, p_v)$ such that $t < s < v$ and $x_t^k = x_v^k = 1$. The last sum counts the number of pixels pairs $(p_t, p_v)$ such that $t < v < s$ and $x_t^k = 1$, $x_v^k = 0$.

Let $C^−$ and $C^+$ be as in (6). Recall that each of them can be computed in one pass over the line. Then the second sum in (11) is simply $C^−(s) \cdot C^+(s)$. For the other two sums, we need additional running sums.

Denote by $A^−(s)$ the number of pixel pairs $(p_t, p_v)$ preceding pixel $p_s$ such that $x_t^k = 1$, $x_v^k = 0$ and pixel $p_t$ precedes pixel $p_v$.

$$A^−(s) = \sum_{(p_t, p_v) \in \Omega \mid t < v < s} (1 - x_v^k) \cdot x_t^k.$$  

Similarly, we define $A^+(s)$ the number of pixel pairs $(p_t, p_v)$ succeeding pixel $p_s$ such that $x_t^k = 0$, $x_v^k = 1$ and pixel $p_t$ precedes pixel $p_v$.

$$A^+(s) = \sum_{(p_t, p_v) \in \Omega \mid s < v < t} (1 - x_t^k) \cdot x_v^k.$$  

Given $C^−$, we compute $A^−$ in one pass over line $l$ using $A^−(0) = A^−(1) = 0$ and recurrence $A^−(s) = A^−(s-1) + (1 - x_{s-1}^k) \cdot C^−(s-1)$.

$A^+$ is computed analogously to $A^−$, given $C^+$. Then the first sum in (11) is $A^+(s)$ and the third sum is $A^−(t)$, and $u^k_{appr}(x_s) = x_s \cdot [A^+(s) - A^−(s)] \cdot C^+(s) + A^−(s)].$

Computing $A^−$ and $A^+$ is linear in the number of pixels in a line. For orientation $d_k$, each pixel appears on one line only. Therefore we can compute $A^−$ and $A^+$ for all lines in $O(mN)$ time, where $m$ is the number of orientations and $N = |\Omega|$. Then the unary term for each pixel is computed in $O(m)$ time. Thus the total time to compute Taylor based unary terms for all pixels is $O(mN)$.

C. Quadratic Approximation of $E_{\text{convexity}}$

Linear approximation derived in section III-A works well for most synthetic and natural images. However, we were able to design synthetic examples (see Fig. 14 and Fig. 15) where linear approximation yields poor results for some initialization. In this section we derive a quadratic approximation that gives better results but at a larger computational cost.

As before, we derive approximation $E_{appr}^k(x)$ for $E_{\text{convexity}}(x)$ in (4) around current solution $x^k$. It is sufficient to derive a quadratic approximation $\phi^k(x_s, x_t, x_v)$ for a clique $\phi(x_s, x_t, x_v)$ and then substitute into (9).

Similarly to linear approximation in Sec. III-A, we first try second-order Taylor expansion of (3) around $x^k$. Omitting constants, we get:

$$\phi^k(x_s, x_t, x_v) = x_s \cdot x_v - x_s \cdot x_v - x_s \cdot x_t \cdot x_v^k + x_s \cdot x_t \cdot x_v^k + x_t \cdot x_v^k + x_v^k \cdot x_t \cdot x_v.$$  

The second and third lines in (12) contain submodular quadratic and unary terms. The first term in (12) is supermodular and therefore cannot be optimized directly using standard techniques. We tried to replace the non-submodular term with two different approximations: (i) $x_s \cdot x_v \approx x_s^k \cdot x_v + x_s \cdot x_v^k$, i.e. Taylor-based unary approximation; (ii) $x_s \cdot x_v \approx 0$, i.e. zero approximation. Both produce poor results (omitted) on the difficult example in Fig. 15.

We now derive an alternative quadratic approximation. Recall from Sec. III-A that linear approximation in (11) can be derived from parallel ICM [16]. First we give details of this derivation and later extend it to quadratic case.

In ICM, given current labeling $x^k$, we fix the labels of all pixels except one, and optimize over the remaining variable. For example, fixing all labels except at pixel $p$, the clique potential in (3) reduces to a unary term $x_s \cdot (1 - x_t^k) \cdot x_v^k$. In parallel ICM this is done simultaneously for all pixels, thus explaining the name parallel. For a given triple clique $(p_s, p_t, p_v)$, summing up the reductions for pixels $p$, $q$ and $r$, and omitting the constant term $x_s^k \cdot x_t^k \cdot x_v^k$, gives exactly the same linear approximation as in Taylor based approximation derived in (10). This is a standard single pixel ICM, which we distinguish from a more general version described below.

The propose a new quadratic approximation based on parallel pairwise (two pixel) ICM. Similarly to single pixel ICM, given current labeling $x^k$, we now fix the labels of all pixels except a pair of interacting pixels. For example, fixing the labels of all pixels expect $(x_s, x_t)$, we get the following quadratic reduction of the clique potential in (3):

$$x_s \cdot x_v^k - x_s \cdot x_t \cdot x_v^k.$$  

Repeating the same procedure for pixel pairs $(q, r)$ and $(p, r)$ and summing up all the reductions results in:

$$\phi^k(x_s, x_t, x_v) = x_s \cdot x_v^k - x_s \cdot x_t \cdot x_v^k + x_s \cdot x_v^k - x_s \cdot x_t \cdot x_v^k + x_s \cdot x_v^k - x_s \cdot x_t \cdot x_v^k + x_s \cdot x_v^k - x_s \cdot x_t \cdot x_v^k.$$  

Note that the approximation in (13) contains a non-submodular term $x_s \cdot x_v$ and, therefore, cannot be efficiently optimized with standard techniques. As previously, we can replace the supermodular term with its linear approximation. We tried two different approximations: $x_s \cdot x_v \approx x_s^k \cdot x_v + x_s \cdot x_v^k$ and
Fig. 4. First row shows synthetic images with added noise $\sigma_{\text{noise}} = 0.2$; Second and third rows show contours and masks of segmentation, $\omega = 0.1$. We used log-likelihood appearance terms, $(\mu_f = 0, \sigma_f = 0.1)$ and $(\mu_b = 1, \sigma_b = 0.1)$. The convexity prior removes noise, connects components and fills holes while preserving sharp corners.

$x_s \cdot x_v \approx 0$, and the second one works better in practice. Thus, our final quadratic approximation based on pairwise ICM is:

$$
\phi_k(x_s, x_t, x_v) = x_s \cdot x_v^k - x_s \cdot x_t \cdot x_v^k
- x_s \cdot x_t^k \cdot x_v
+ x_s^k \cdot x_v - x_s^k \cdot x_t \cdot x_v
$$

Observe that pairwise ICM approximation in (14) is different from the second order Taylor approximation in (12) even though the single pixel ICM approximation coincides with the first order Taylor approximation.

Pairwise ICM approximation is more accurate than linear approximation. It coincides with the original triple clique potential $\phi(x)$ on at least five out of eight configurations. More precisely depending on the current configuration $x^k$ the approximation coincides on either five, six or seven configurations. This is in contrast to at least four out of eight configurations for the linear approximation. Pairwise ICM approximation (14) works better than linear approximation, but is less efficient (see Sec. IV-D3). In each iteration, two new submodular pairwise cliques are created for each triple clique based on current solution. Furthermore, while dynamic programming can be used to efficiently compute unary terms, the same ideas do not apply to pairwise terms. Thus, optimization of quadratic approximation is more time consuming, compared to the unary case. Nonetheless, approximation based on pairwise ICM may be useful for optimization of energies other than $E_{\text{convexity}}$.

IV. EXPERIMENTS

Below we apply convexity shape prior to image segmentation. We discretized orientations using $11 \times 11$ stencil yielding 40 orientations for all synthetic images. For natural images we found a $5 \times 5$ stencil yielding 8 orientations sufficient. Note, since the convexity term is NP-hard and our optimization is local, in general, we cannot guarantee convex results. However in practice the number of violations is either zero or negligible.

We collected a new dataset with ground truth convex segmentations. The code and the dataset are available from http://vision.csd.uwo.ca/code/ and http://vision.csd.uwo.ca/code/.

The remainder of this section is structured as follows. First in Sec.IV-A we apply our method as described in Algorithm 1 (i.e. using linear approximation and dynamic programming) to synthetic and natural images. We call this approach “Direct” to distinguish it from subsequent variants. Next, in Sec.IV-B we experiment with alternative central-clique and local convexity models. In Sec. IV-C we compare our “Direct” optimization with existing state-of-the-art optimization methods. Finally, in Sec. IV-D we discuss limitations of our direct approach and evaluate two possible solutions: Gradual Non-Submodularization in Sec. IV-D2 and trust region based on quadratic approximation in Sec. IV-D3.
A. Direct approach: Trust Region with Linear Approximation

First we validate our method on synthetic images with noise $N(0,0.2)$, see Fig. 4. We assume given target appearance distributions for foreground and background, and combine standard log-likelihood data terms with the convexity shape prior

$$E(x) = E_{appearance}(x) + E_{convexity}(x).$$

Here $E_{appearance}(x) = \sum_{p \in \Omega} D_p(x_s)$ is the appearance term, $D_p(x_s) = -\log Pr(I_p|x_s)$ and $E_{convexity}(x)$ is as in (4). Fig. 4 demonstrates that our convexity prior removes noise, insures connectivity and fills in holes while preserving sharp corners.

Next, we use convexity prior in interactive segmentation of natural images with user scribbles. The convexity prior is especially useful for images where there is an overlap between the foreground and background appearance, see Figures 6-10. Such overlap often leads to holes in the foreground or larger parts of the background erroneously segmented as the foreground. The convexity prior prevents such results. Length regularization is either too weak to remove the noise or too strong causing shrinking.

We now specify the details of our interactive segmentation. For appearance we use the recent submodular $L_1$ color separation term proposed in [26]. This term is based on $L_1$ distance between unnormalized histograms of foreground and background colors. Unlike standard appearance models, the color separation does not require re-estimation of the parameters and can be efficiently and globally optimized. We use 16 bins per color channel and combine the color separation term with the convexity prior, subject to hard constraints on the user scribbles

$$E(x) = E_{L1}(x) + E_{convexity}(x).$$

We then compare with the standard length regularization

$$E(x) = E_{L1}(x) + \sum_{(p,q) \in \mathcal{N}} \omega[x_s \neq x_t].$$

Figures 6-8 show segmentation results on two natural and one medical image. We vary the weight $\omega$ for our convexity prior in (3) and optimize as discussed in Sec. III. Similarly, we vary the weight $\omega$ for the length regularization and optimize with one graph-cut [26]. We show the resulting segmentations and compare them with the results obtained without regularization. We use green frames - for convexity, blue - for length. For length regularization we chose values $\omega$ for which the segmentations change significantly. The length regularization is either too weak to remove the noise or too strong and has a shrinking bias. Figure 10 shows similar results on additional natural images demonstrating the sensitivity of the length regularization and robustness of convexity shape prior.

We experiment both with contrast sensitive length ($\omega$ depends on pixel pair $(p,q)$ as in [4]) and pure length, see Fig. 7. There is no significant difference in their performance. The same sensitivity to the parameter $\omega$ is observed; compare the red (contrast sensitive) frames with the blue ones (pure length).

Our model is parameter-free due to infinity cost constraints. In practice, we have to choose finite $\omega$. There is almost no
variation in segmentation results for different values of $\omega$, once the value is high enough, making it a very robust regularizer. In fact, for each image we can compute finite $\omega$ such that violating a single constraint is more expensive than the initial solution. In cases where using such large $\omega$ leads to poor local minimum, gradually increasing the weight $\omega$ (annealing) can potentially escape to a better solution (see Gradual Non-Submodularization in sec. IV-D2).

There is also very little variation in segmentation results with change of scale. Figure 9 shows segmentation results obtained on several natural images and their 25% and 11% scaled versions using the same parameters for $\omega$ and appearance. User scribbles were scaled accordingly in each experiment.

**B. Alternative convexity models**

Below we discuss some variations of our convexity model.

1) Central Cliques Model: One natural question regarding our model is whether we need all the triple cliques on a line to enforce convexity. Indeed, it is sufficient to use a smaller subset consisting only of central triple cliques $(p, q, r)$, i.e., $|p - q| = |q - r|$, see example on the right. This reduces the number of triple cliques from $O(n^3)$ to $O(n^2)$ for a line with $n$ pixels. However, our dynamic programming procedures for evaluation (Sec. II-A) and for approximation (Sec. III-B) are no longer applicable. Brute force computation takes $O(n^2)$ operations per line with $n$ pixels, as opposed to linear time with our dynamic programming for the full model. Nonetheless, we compare between our full model and the central cliques model, see Fig.11. Since the resulting segmentations have no convexity violations their energies can be directly compared. The energy is slightly better with the central cliques, but its running time is 25-30 times longer. The difference in time will be even more significant for larger images.

2) Local Convexity Model: Another natural variation of our model is the local convexity model, in which we penalize local spatially symmetric configurations 1-0-1 within a fixed size neighborhood around each pixel. The number of triple cliques is linear in the number of image pixels and the neighborhood size. Similarly to our full model, the local convexity model can be optimized using our trust region method. Our dynamic programming is not directly applicable. The local model can only enforce local convexity. Namely, it is blind to violations of convexity that lie at a scale larger than the neighborhood size. Therefore the results can contain holes, multiple connected components and large scale concavities. Consider a $100 \times 100$ synthetic image example in Fig. 13. With a small neighborhood size, the number of triple cliques is small and computation is efficient, but the results are far from convex (Fig. 13, top-right). Moreover, small neighborhood size yields very coarse discretization of orientation. As we increase the neighborhood size, the number of convexity violations is reduced (Fig. 13, bottom row), eventually reaching a convex segmentation for a large enough neighborhood. However, the...
Input User Scribbles Length Regularization Convexity Regularization
\(\omega=0.005\) \(\omega=1\) \(\omega=0.32\) \(\omega=0.33\) \\
\(\omega=0.005\) \(\omega=1\) \(\omega=0.795\) \(\omega=0.796\) \\
\(\omega=0.005\) \(\omega=1\) \(\omega=2\) \(\omega=8\) \\
\(\omega=0.005\) \(\omega=0.1\) \(\omega=0.57\) \(\omega=0.575\)

Fig. 10. Additional results comparing length and convexity regularizers. Except for the second row, all images demonstrate sensitivity to length weight \(\omega\).

\[E_1=15020, T=4.6 \text{ sec.} \quad E_2=14666, T=122 \text{ sec.}\]

Fig. 11. Comparison between the full convexity model with all triple cliques vs. central cliques convexity model. The later does not allow efficient evaluation and approximation of \(E_{\text{convexity}}\) using dynamic programming and therefore is much slower.

model becomes very slow to optimize.

C. Alternative Optimization Methods

The most related optimization method to ours is LSA [9], which is also based on trust region framework. However, it was designed for non-submodular energies with only pairwise cliques. For this class of energies, LSA reports state-of-the-art results [9]. LSA approximates the energy by replacing non-submodular pairwise cliques with their Taylor expansions while preserving all submodular pairwise cliques. Even though our convexity prior is not pairwise, it is possible to reduce each triple clique to several pairwise potentials using an additional auxiliary node [15] and optimize them with LSA. We call this reduced version r-LSA. Reducing all triple cliques would result in a prohibitively large number of auxiliary nodes and pairwise cliques for our full convexity model. Even for the central clique model the number of resulting pairwise cliques is quite large. For an \(n \times n\) image central cliques model with \(m\) orientations produces \(O(mn^3)\) pairwise potentials, which is very costly both to optimize and to evaluate. Nonetheless, we tried this approach on a small \(91 \times 122\) image. The first two rows in Figure 12 compare r-LSA approach to our method.

We apply both methods to the central clique model\(^\text{1}\) and vary \(\omega\). Note that r-LSA is an order of magnitude slower than the slow version of our method. As the value of \(\omega\) increases, r-LSA fails to obtain satisfactory solutions. We believe that there could be serious drawbacks in splitting clique \(\phi(x_s, x_t, x_v)\) into individual submodular and supermodular parts and then approximating the supermodular part. One sign of a problem is that there are infinitely many such decompositions and it is not clear which one would give a better approximation.

Our full model with all triple cliques is also prohibitively expensive for standard optimization methods designed for non-submodular energies, such as QPBO [2] and TRWS [14]. However, we can use these methods to optimize the more compact central clique model as well. The last two rows in Figure 12 show segmentation results of QPBO and TRWS for several values of \(\omega\). For values of \(\omega\) that are not sufficiently large to enforce convexity, all four methods, QPBO, TRWS, r-LSA and Trust Region, return globally optimum, but useless solutions. However, when \(\omega\) is large enough, QPBO, TRWS

\(^{1}\)Even though the full model is more efficient for our method, for this experiment we use central cliques to have identical energies for direct comparison.
Fig. 12. Comparison between our method without dynamic programming (no DP), r-LSA, QPBO and TRWS on the central clique model with 8 orientations. We use thresholded appearance terms for initialization when needed. As $\omega$ increases, making the energy more difficult, QPBO was not able to label any pixel (shown in gray) and TRWS did not converge after 5000 iterations, which took several hours. For $\omega$ large enough to enforce convexity, all methods except ours fail.

Fig. 13. Local Convexity Model: penalizes local 1-0-1 configurations within a fixed neighborhood around each pixel. Small neighborhoods are efficient to optimize but have very coarse discretization of orientation. Moreover, they are sensitive to noise and might result in multiple disconnected components. Larger neighborhood are slow to optimize.

and r-LSA fail to obtain a satisfactory result. Our trust region approach obtains good results in all cases.

D. Optimization Limitations and Proposed Solutions

1) Local Minimum: Trust region framework is a local iterative optimization and therefore we can only guarantee a local minimum [10]. Figure 14 demonstrates some sensitivity with respect to initialization. A trivial initialization with all pixels in the foreground, denoted by “init 1” and delineated by the red contour, leads to a local minimum. Initializing with the maximum likelihood label per pixel, denoted by “init 2” results in solution with better energy. Empirically, we obtain better results starting with maximum likelihood labels. This is consistent for all the experiments, both on synthetic and real images.

2) Gradual Non-Submodularization: As shown in the experiments with the natural images, segmentation with convexity shape prior yields robust results for different values of the weight $\omega$, once they are high enough. However, we could design a synthetic example (Fig. 15) where caution should be used when selecting the weight $\omega$ in (3). High values of $\omega$ tend to cause more aggressive optimization steps, and might lead to a solution that is convex but far from a global optimum (Fig. 15, top row, third column). Using low values of $\omega$ allows deviation from convexity (Fig. 15, bottom-left). To obtain a convex solution that avoids such local minima, the weight $\omega$ of the convexity shape prior can be increased gradually, similarly to the idea of “Graduated Non-Convexity” [1]. We can start with a very low weight, where the energy is mostly guided by the submodular appearance terms and is easier to optimize. Then the result is used for initialization of the next round, in which the weight $\omega$ is slightly increased. See an example of such gradual non-submodularization on a synthetic image in Fig.15 (bottom row).

For our discrete energy, gradual increase in weight $\omega$ makes the energy gradually more non-submodular, more non-linear and, therefore, more difficult. Similar improvements over
direct aggressive optimization are obtained in other synthetic (see Fig. 14) and medical (see Fig. 16) images.

In Fig. 16 we apply convexity shape prior for liver segmentation on an MRI image. Here again, high values of \( \omega \) result in a convex but not accurate solution (top-right). Values of \( \omega \) that are too low allow deviation from convexity (bottom-left). **Gradual non-submodularization** (bottom row) escapes such bad local minima and results in a solution with a lower energy.

Gradual non-submodularization can also alleviate the sensitivity of our trust region method to initialization. In Fig. 14 we were able to obtain the same global optimum starting with either “init 1” or “init 2” for both synthetic examples.

To further evaluate the advantage of gradual non-submodularization we collected a database of 51 natural images. We manually segmented each image to obtain ground truth and convexified the resulting foreground segment to ensure that ground truth is convex. We performed two sets of experiments: using 5\( \times \)5 orientations stencil (i.e. 8 orientations) and 11\( \times \)11 stencil (i.e. 40 orientations.) In each experiment we run both, our direct approach with \( \omega = 10 \) and the gradual non-submodularization approach with the schedule \( \omega = \{0.0001, 0.001, 0.01, 0.1, 1, 10\} \). All methods were initialized with the solution that minimizes submodular \( L_1 \) color separation term proposed in [26] subject to user scribbles constraint. Recall that \( L_1 \) color separation does not require re-estimation of the parameters and can be efficiently and globally optimized. We used 16 bins per color channel.

We then compared the results in terms of mean running time (in sec.), distance from the ground truth (percentage of misclassified pixels) and energy (percentage of images in which gradual approach obtains strictly lower energy. Table I summarizes the results. While having a slightly longer mean running time, gradual approach achieves more accurate results with respect to the ground truth for both 5\( \times \)5 and 11\( \times \)11. In terms of energy, gradual approach achieves lower energy in more than 84\% of the images.

Figure 17 shows several images from the database. The database can be downloaded from http://vision.csd.uwo.ca/data/. Rows 1, 4-6 show examples where gradual method obtains lower energy than the direct approach with both 5\( \times \)5 and 11\( \times \)11 orientation stencils. Second row shows an example where direct and gradual approach converge to the same solution. Third row shows an example where in case of 11\( \times \)11

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### Table I

<table>
<thead>
<tr>
<th>Method</th>
<th>Energy (%)</th>
<th>Distance (%)</th>
<th>Running Time (sec.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direct</td>
<td>6075</td>
<td>4975</td>
<td>529</td>
</tr>
<tr>
<td>Gradual Non-Sub</td>
<td>4275</td>
<td>8150</td>
<td>477</td>
</tr>
<tr>
<td>Submodularization</td>
<td>17475</td>
<td>8150</td>
<td>11</td>
</tr>
<tr>
<td>Starting with Init 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Starting with Init 2</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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**Fig. 14.** Local optimization of convexity shape prior might yield different segmentation results for different initializations. We used an 11\( \times \)11 mask to discretize orientation, \( \omega = 100 \) and given target appearance models \( \mu_{fg} = 0, \sigma_{fg} = 0.1 \). \( \mu_{bg} = 1, \sigma_{bg} = 0.1 \). Gradual Non-Submodularization yields global minimum solutions (verified by geometrical arguments) using any initialization (see text for details).

**Fig. 15.** Gradual Non-Submodularization with Linear Approximation vs. Quadratic Approximation. First row: input image, initial solution based on data terms, direct optimization with linear approximation resulting in weak local minimum, and direction optimization with quadratic approximation resulting in a satisfying solution. Second row: Gradual non-submodularization. Each increase in \( \omega \) starts new optimization using previous result as initialization. We used standard log-likelihood data terms: \( \mu_{fg} = 0, \sigma_{fg} = 0.1 \). \( \mu_{bg} = 1, \sigma_{bg} = 0.1 \).
The slow running time we chose very small $50 \times 50$ yet challenging synthetic binary images with artificially added noise. In nine out of ten images, quadratic approximation yielded solutions with lower energy. The average running time was 27.15 and 36.75 sec. for linear and quadratic approximation models respectively. Note that the small difference in running time is due to the small size of the images. Figure 18 shows input images (leftmost column), initial solutions (second column), results obtained with linear approximation (third column) and results obtained with quadratic approximation (rightmost column). Though for our dense convexity model quadratic approximation is very inefficient, it might be useful for other high-order energies where the number of high-order cliques is not as high.

V. CONCLUSION AND FUTURE WORK

We propose convexity prior as a new regularizer and develop efficient discrete optimization based on trust region and dynamic programming. Our convexity prior does not have shrinking bias and is robust w.r.t. parameter $\omega$.

In the future, we plan to explore meaningful extensions of strict convexity. For example, we can explore contrast sensitive convexity, which is similar to contrast sensitive length. The penalty for convexity violation by triplet $(p_x, p_t, p_v)$ can carry a smaller weight if there is a high contrast on a line connecting pixels $p_x$ and $p_v$. This formulation is straightforward, but the main difficulty will be extending our dynamic programming algorithms to handle this case. Another direction is to extend our model to handle objects with multiple convex parts.

VI. ACKNOWLEDGEMENTS

We are thankful for generous support by Canadian NSERC Discovery and RTI Programs.

REFERENCES

Fig. 17. Qualitative comparison of direct and gradual approach using $5 \times 5$ and $11 \times 11$ orientation stencils. See text for details.

Fig. 18. Qualitative comparison of linear and quadratic approximation using $11 \times 11$ orientation stencils. Left-to-right: Input image, initial solution using maximum likelihood label per pixel based on the appearance terms, solution with linear approximation, solution with quadratic approximation. We used standard log-likelihood data terms: $(\mu_{fg} = 0, \sigma_{fg} = 0.4)$, $(\mu_{bg} = 1, \sigma_{bg} = 0.4)$. Quadratic approximation obtained lower energy for 9 out of 10 images. Green frame highlights the negative example.


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