Centroidal Power Diagrams with Capacity Constraints: Computation, Applications, and Extension

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Abstract

This article presents a new method to optimally partition a geometric domain with capacity constraints on the partitioned regions. It is an important problem in many fields, ranging from engineering to economics. It is known that a capacity-constrained partition can be obtained as a *power diagram* with the squared L2 metric. We present a method with super-linear convergence for computing optimal partition with capacity constraints that outperforms the state-of-the-art in an order of magnitude. We demonstrate the efficiency of our method in the context of three different applications in computer graphics and geometric processing: displacement interpolation of function distribution, blue-noise point sampling, and optimal convex decomposition of 2D domains. Furthermore, the proposed method is extended to capacity-constrained optimal partition with respect to general cost functions beyond the squared Euclidean distance.

Keywords: centroidal power diagram, displacement interpolation, convex decomposition, blue noise

$\textbf{Concepts: } \bullet \textbf{Computing methodologies} \rightarrow \textbf{Computer graphics;}$

1 Introduction

The central theme of the present paper is to apply and extend the notion of power diagram to computing optimal partition of a geometric domain with capacity constraints. A partition of a geometric domain is a collection of disjoint constituent parts whose union is equal to the whole domain. A partition is said to be optimal if all its partitioned regions, also called *cells*, have compact shapes with respect to some specified cost *kernel* that defines the cost function. Here "compactness" can be roughly measured by the isoperimetric quotient. Capacity constraints on a partition mean that each cell of the partition has a specified size, measured in volume (or area in 2D) or mass. Hence, an optimal partition with capacity constraints is a partition such that its cells have compact shapes as well as sizes

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equal to some specified quantities, unlike the optimal mass transport [Santambrogio 2015] that concentrates on capacity constraints regardless of shape compactness or the centroidal Voronoi tessellation [Du et al. 1999; Liu et al. 2009] that considers the opposite.

Finding an optimal partition by some criteria is a central issue in many applications. Optimal partitions, with or without capacity constraints, are used in computer graphics and geometric computing for point sampling [de Goes et al. 2012], information visualization [Balzer et al. 2005], shape segmentation [Lu et al. 2007], shape matching [Su et al. 2015], area-preservation mapping [Zhao et al. 2013] and mesh generation [Du et al. 1999; Liu et al. 2009]. The optimal partitioning problem with capacity constraints is also studied in engineering and economics for optimal resource location or provision of services. For example, it appears as the optimal coverage problem in robotics and sensor networks [Cortés 2010; Patel et al. 2014], the pattern formation problem in material science [Bourne et al. 2014], and the land consolidation problem in agriculture [Borgwardt et al. 2014].

The contribution of this paper is twofold.

- An L-BFGS method with empirically superlinear convergence is developed for computing capacity-constrained centroidal power diagram, assuming the squared Euclidean distance as the cost kernel. We show its superiority to previous method and validate it in two applications: displacement interpolation of function distribution and blue-noise point sampling. Furthermore, the method is extended to computing optimal capacity-constrained partition with a general cost function;
- 2. Based on the centroidal power diagram, an effective method is developed for decomposing a 2D polygonal region into convex and compact cells with capacity constraints.

2 Related Work

Voronoi Diagram & Power Diagram The Voronoi diagram is a partition of a given domain $\Omega \subset E^d$ into regions based on the straight-line distance to a set of points $\{x_i \in \Omega\}_{i=1}^n$ (called *sites* or *generators*), where x_i dominates the subregion

$$\Omega_i: \{x \in \Omega \mid ||x - x_i|| \le ||x - x_j||, j \ne i\},\$$

which is called the *Voronoi cell* of the site x_i . A power diagram is defined as an extension of Voronoi diagrams, where each site x_i has an associated coefficient (called a "weight") and dominates the subregion

$$\Omega_i: \{x \in \Omega \mid ||x - x_i||^2 - w_i \le ||x - x_j||^2 - w_j, j \ne i\}.$$

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 Ω_i is called the *power cell* of the site x_i . The power diagram reduces to the Voronoi diagram when all the weights are equal [Aurenhammer 1987; Okabe et al. 1992].

Centroidal Voronoi Tessellation (CVT) A centroidal Voronoi tessellation (CVT) of a set of sites $X = \{x_i\}_{i=1}^n$ is a special Voronoi diagram for which each site x_i coincides with the centroid (i.e. the center of mass) of its Voronoi cell Ω_i [Du et al. 1999]. Alternatively, a CVT can be also defined as a stationary point of the following objective function:

$$Q(X) = \sum_{i=1}^{n} \int_{\Omega_i} ||x - x_i||^2 \rho(x) \mathrm{d}\sigma, \qquad (1)$$

where $\rho(x)$ is a C^1 -smooth density function on the whole domain Ω . Domain partition using CVT yields convex polyhedral cells (barring those abutting boundaries) that have compact shapes and similar sizes, assuming that $\rho(\cdot)$ is uniform. Due to these properties, CVT is widely used for computing optimal partitions in various applications [Lu et al. 2014; Zhou et al. 2015], and in some cases even for optimal partition with capacity constraints [Balzer and Heck 2008; Chen et al. 2012; Li et al. 2010]. However, it is known that the class of partitions given by the Voronoi diagram is too restrictive to properly accommodate capacity constraints [Zhang et al. 2016]. A prevailing method for computing CVT is Lloyd's method, which behaves as a gradient-descent method and thus converges linearly. A quasi-Newton method for computing CVT is proposed in [Liu et al. 2009] that is based on the limited-memory BFGS technique and is shown to demonstrate super-linear convergence in practice.

Optimal Mass Transport (OMT) The optimal mass transport (OMT) was formalized by the French mathematician Gaspard Monge [1781] who studied the most economical way of moving soil from one area to the other and introduced the theory in terms of minimizing the L_1 norm of the distance transported. The problem of minimizing L_2 cost whose kernel is the squared distance (note that the square root of the minimum L_2 cost is called the 2-Wasserstein metric), however, takes on additional structure and can be expressed as the fully nonlinear degenerate elliptic Monge-Ampère equation. In 1949, Kantorovich proved the existence and uniqueness of the optimal transport plan and showed how Monge's problem is connected to linear programming [Kantorovich and Gavurin 1949]. The Monge-Kantorovich optimization technique has been studied in a wide range of research fields in the past two decades, including imaging, adaptive meshing, geophysical fluid dynamics, and cosmology.

It is shown in [Pogorelov 1994], [Gangbo and Cann 1996] (example 1.6) and [Aurenhammer et al. 1998] (or as a direct consequence of [Brenier 1991]) that for the semi-discrete OMT (from a discrete probability distribution function to a continuous one), the class of partitions parameterized by the power diagram provides the proper representation to accommodate capacity constraints in the following sense. Given a set of fixed sites in a domain, there exists a unique set of weights (up to an additive constant) such that the power cells, which are determined by the given sites and the associated weights, satisfy the specified capacity constraints and minimize the sum of least squares costs of all the cells. Here, the least squares cost of the cell Ω_i associated with a site x_i is defined to be $\int_{\Omega_i} \|x - x_i\|^2 \rho(x) d\sigma$. This means that we just need to look for the capacity-constrained partition within the set of power diagrams with different weighting schemes, rather than having to search in the set of arbitrary partitions of the domain, which obviously cannot be parameterized by a finite number of parameters.

Centroidal Power Diagram (CPD) In this paper, we are interested in computing the optimal partition of a geometric domain with specified capacity constraints. Similar to the conventional OMT problem, the solution to such a capacity-constrained optimal partition problem with the least squares cost function can be represented by a power diagram [Balzer et al. 2009]. More specially, the solution is given by a centroidal power diagram (CPD) which is a special power diagram in which each site coincides with the centroid of its power cell. Interestingly, the weights of the power diagram exactly correspond to the degrees of freedom required to account for capacity constraints. However, one should notice that the relation between the weights and the capacities of the cells is non-trivial (in this context, the term "weight" is misleading, they are completely different from the masses of the cells).

CPD was used for blue-noise sampling in computer graphics. In numerical implementation, Balzer et al. [2009] proposed several iterative schemes to compute capacity-constrained optimal partitions, including the method based on pixel-based clustering and the methods that enforce capacity constraints on the Voronoi diagram as well as the power diagram. These methods are easy to implement. But because they update only one single pixel or site in each iteration, the methods have very slow convergence and low efficiency. Later, a Lloyd-like method that repeatedly moves each site to the mass center of its cell is proposed to compute CPD for blue-noise sampling [de Goes et al. 2012], which has only linear convergence.

The above discussions naturally lead to two questions. First, is it possible to develop an efficient method that outperforms these existing methods? Second, remarking that the least squares cost function is in fact the squared L_2 metric, we want to know whether the new numerical method is able to deal with the CPD problem with a general cost. We will give positive answers to both questions in this paper.

3 Capacity-Constrained Optimal Partition

3.1 Problem Formulation

We first recall the notions used for a power diagram in E^d , ddimensional Euclidean space. Let $\Omega \subset E^d$ be a convex, closed, bounded and connected domain. Let $X = \{x_i\}_{i=1}^n \subset \Omega$ be a set of points, called *sites*, with its associated *weights* $\{w_i\}_{i=1}^n \subset R$. Then the power diagram induced by the weighted points (x_i, w_i) is the collection of convex polytopes, or *cells*, $PD = \{PD_i\}_{i=1}^n$, where $PD_i = \{x | ||x - x_i||^2 - w_i \leq ||x - x_j||^2 - w_j, \forall j, j \neq i\}$. The restriction of the power diagram PD to the domain Ω induces a partition of Ω , denoted by $PD_\Omega = \{PD_i \cap \Omega\}_{i=1}^n$.

Now consider partitioning a domain Ω in Euclidean space E^d into a set of n regions $R = \{R_i\}_{i=1}^n$ in a capacity-constrained manner. Let $X = \{x_i\}_{i=1}^n \subset \Omega$ denote n given points, also called *sites*, with associated capacity constraints $c_i > 0, i = 1, 2, ..., n$. We define the least squares cost of each region R_i , given by x_i , to be $\int_{R_i} ||x - x_i||^2 \rho(x) d\sigma$, where $\rho(x)$ is a C^1 -smooth density function on Ω . The capacity constraint here requires that $|R_i| \equiv \int_{R_i} \rho(x) d\sigma = c_i, i = 1, 2, ..., n$, that is, the capacity (i.e. area or volume) of each region R_i matches the constraint c_i . (Of course, $\sum_i c_i$ is assumed to be equal to $|\Omega| \equiv \int_{\Omega} \rho(x) d\sigma$.) Then the problem of *capacity-constrained optimal partition* is to find the partitions R_i and the sites x_i such that the total cost

$$Q(X,R) = \sum_{i=1}^{n} \int_{R_i} ||x - x_i||^2 \rho(x) d\sigma$$
 (2)

is minimized, subject to the constraints $|R_i| = c_i, i = 1, 2, ..., n$.

In the case of the L2 cost, it follows from [Aurenhammer et al. 1998] that the solution $(X^*, R^*) = \arg \min Q(X, R)$ exists and is given by a power diagram; See [Pogorelov 1994] and [Gangbo and Cann 1996] for a proof, as well as the derivations in [Gu et al. 2013]. Aurenhammer [1998] further showed that the optimal power diagram can be found by extremizing

$$F(X,W) = \sum_{i=1}^{n} \int_{PD(W)_{i}} ||x - x_{i}||^{2} \rho(x) \mathrm{d}\sigma - \sum_{i=1}^{n} w_{i}(|PD(W)_{i}| - c_{i})$$
(3)

where $PD(W)_i$ (or PD_i) is a power cell dominated by x_i . A key observation made in [Aurenhammer et al. 1998] is that, when the sites X in F(X, W) are fixed, their weights can be computed as a maximizer of a concave function to make the cells of the resulting power diagram meet the capacity constraints $|PD_i| = c_i$, i = 1, 2, ..., n. This observation leads to a Lloyd-like method [Cortés 2010; de Goes et al. 2012] for minimizing the objective function F(X, W) in Eqn. (3), i.e., alternatively optimizing the weights to meet the given capacity constraints and moving the sites to their respective mass centers. We will discuss these numerical techniques in detail in the next section, and introduce a new method that better exploits the derivatives of the objective function, resulting in a faster convergence.

3.2 Partition with General Cost Functions

Given a general distance cost kernel d(x, y), where $x, y \in \Omega \subset E^d$, a generalized power diagram (also called a Laguerre diagram) of Ω with the weighted sites $\{(x_i, w_i)\}_{i=1}^n$ is defined to be the collection of cells $GD = \{GD_i\}_{i=1}^n$, where $GD_i = \{x|d(x, x_i) - w_i \leq d(x, x_j) - w_j, \forall j, j \neq i\}$. When d(x, y) is the squared distance, the generalized power diagram reduces to the traditional power diagram whose cells are convex polytopes. However, with a general cost, the cells GD_i are not necessarily convex and their boundaries are curves or curved surfaces rather than straight lines or planes.

Let μ denote a measure with non-vanishing density $\rho(.)$, supported by the domain Ω . Suppose that each site x_i has capacity constraint $c_i > 0$, such that $\sum_i^n c_i = |\Omega| \equiv \int_{\Omega} d\mu$, which is the total mass of the measure μ . Again, let d(x, y) denote a distance cost between any two points x and y in Ω . Let $R = \{R_i\}_{i=1}^n$ denote a partition of Ω , that is, $\bigcup R_i = \Omega$ and $R_i \cap R_j = \emptyset, \forall i \neq j$. Define the cost of the partition R with respect to the sites $X = \{x_i\}_{i=1}^n$ as

$$Q(X,R) = \sum_{i=1}^{n} \int_{R_i} d(x,x_i) d\mu.$$
 (4)

In what follows, we support that the distance d(x, y) satisfies the twist condition, i.e. $\nabla_x d(x_0, y)$ is one-to-one $\forall x_0$, which ensures the existence and uniqueness of the optimal transport plan (see e.g. [McCann and Guillen 2010] p. 10 and [Santambrogio 2015]). Then the capacity-constrained optimal partition problem is to find the sites X^* and partition R^* that minimize the function Q(X, R) subject to the *capacity constraints* $c_i = |R_i| \equiv \int_{R_i} d\mu$, $i = 1, 2, \ldots, n$. The difference between this definition of the general setting and the previous simple setting of the squared L_2 cost is that here Ω may be a curved surface and that d(x, y) is a general cost kernel.

Next we shall show that the optimal partition R^* with capacity constraints assuming a general cost function Q(X, R)) can be found by only considering the candidates of extended power diagrams where each region (possibly with curved boundaries) is associated with a weight. This is a natural extension from [Aurenhammer et al. 1998; Santambrogio 2015]. Its proof is given in the appendix. **Theorem 1:** For any fixed sites $X = {x_i}_{i=1}^n$, the capacityconstrained partition $R = {R_i}_{i=1}^n$ of the domain Ω that minimizes the cost function in Eqn. (4) is the one such that the boundary of any two adjacent regions R_a and R_b , which are associated with the sites a and b in X respectively, is defined by the equation $d(x, a) - w_a = d(x, b) - w_a$ for some constants w_a and w_b .

Similar to the treatment in [Aurenhammer et al. 1998] for the case of the least squares cost function, we build a new objective function to accommodate the optimal partition according to the Lagrangian Multiplier Theorem:

$$F(X,W) = \sum_{i=1}^{n} \int_{GD(W)_{i}} d(x,x_{i}) d\mu - \sum_{i=1}^{n} w_{i}(|GD(W)_{i}| - c_{i}).$$
(5)

It can be shown that F(X, W) is concave w.r.t. W and therefore has a unique maximizer (up to an additional constant), denoted $W^* = \{w_i^*\}_{i=1}^n$. When the sites $X = \{x_i\}_{i=1}^n$ are fixed, the optimal partition exists uniquely and is given by the generalized Voronoi diagram $GD(W^*)$. When the capacity constraints are met, F(X, W) is exactly equal to the cost function in Eqn. (4). Furthermore, the gradient of F(X, W) w.r.t. W can be shown to be [Aurenhammer et al. 1998; Santambrogio 2015]

$$\nabla_W F(X, W) = -(|GD(W)_1| - c_1, \dots, |GD(W)_n| - c_n),$$
(6)

which is a normal vector to the tangent plane of F(X, W). Hence, to seek the maximizer W^* of F(X, W), we have $\nabla_W F(X, W^*) = 0$, which means that all the capacity constraints are satisfied.



Figure 1: An optimal partition $(X^*, W^*(X^*))$ is a saddle point of the cost function F(X, W) in Eqn. (5).

To summarize, we optimize over all the sites X and weights Wto find the generalized weight Voronoi diagram that gives an optimal partition with capacity constraints. Because there is a unique set of weights W (up to an additional constant) for every set of sites X to make the resulting weighted Voronoi diagram satisfy the capacity constraints, all the capacity-constrained partitions can be represented by a surface $W = W^*(X)$, which will be called the constraint surface (see the green curve in Fig. 1(b)). For each point $(X_a, W^*(X_a))$ on the constraint surface, the restriction $F(X_a, W)$ of F(X, W) (Eqn. (5)) to the subspace $X = X_a$ attains its unique maximum at the point $(X_a, W^*(X_a))$ among all the weighting schemes W (see the red curve in Fig. 1(b)). Meanwhile, the restriction $F(X, W^*(X))$ of F(X, W) to the constraint surface attains its local minimum at a point $(X^*, W^*(X^*))$ that gives an optimal capacity-constrained partition, which is marked with the blue ball on the green curve in Fig. 1(b). Hence, an optimal partition $(X^*, W^*(X^*))$ is a saddle point of the cost function F(X, W) in Eqn. (5).

4 Algorithm for Computing Centroidal Power Diagrams

In this section we propose an efficient method for computing CPDs, as well as the implementation details.

4.1 Overall Algorithm

As discussed above, if (X^*, W^*) defines the optimal partition, the cost function F(X, W) (see Eqn. (5)) is minimized by X^* while maximized by W^* . In other words, finding (X^*, W^*) is neither a pure maximization problem nor a minimization problem. de Goes et al. [2012] suggested updating X and W in an alternative style, where the two key operations are: (1) finding the optimal weighting scheme W to meet the given capacity constraints using Newton's method while fixing X [Benamou et al. 2014]; and (2) moving each site x_i to the mass center of its corresponding power cell. However, this iterative scheme has only linear convergence due to the ineffectiveness of the latter operation. As de Goes et al. [2012] observed, developing an efficient CPD algorithm with super-linear convergence is non-trivial.

Based on the discussion in Section 3, the optimal weights W^* maximizing the cost function in Eqn. (5) is uniquely determined (up to an additional constant) for a given site collection X. Therefore the cost function can be reconsidered as a function of X :

Minimize
$$F(X, W^*(X)) = \sum_{i=1}^n \int_{GD(W^*)_i} d(x, x_i) d\mu$$

 $-\sum_{i=1}^n w_i^* (|GD(W^*)_i| - c_i),$ (7)

where W^* depends on X so that the specified capacity constraints are met. Based on this formulation, we will use the combination of the reduced-gradient technique and the L-BFGS solver [Lasdon et al. 1974; Liu and Nocedal 1989] to move (X, W) on the constraint surface $W = W^*(X)$ towards the minimum of $F(X, W^*(X))$.

Our method works as follows. We start from a feasible point (X_0, W_0) , i.e. a point on the constraint surface $W_0 = W^*(X_0)$. Without loss of generality, we assume that (X_k, W_k) , the position in the k-th iteration, is a feasible point as well. Next, we compute the reduced-gradient vector $\nabla_{x_i} F(X, W^*(X))$ and feed it to the LBFGS solver to update the sites X_k to X_{k+1} . Note that the weights are computed according to the up-to-date sites before we evaluate the objective function F. Thus the (k + 1)-th iteration yields the next position (X_{k+1}, W_{k+1}) . Repeat the process until some stopping criteria is met.

We now consider the derivation of $\nabla_{x_i} F(X, W^*(X))$. Based on the envelope theorem [Milgrom and Segal 2002], we know

$$\nabla_{x_i} F(X, W^*(X)) = \nabla_{x_i} F(X, W).$$
(8)

According to Reynolds' transport theorem [Leal 2007] that recasts derivatives of integrated quantities, we can further show that when the cost kernel d is differentiable everywhere in the domain, the gradients of $F(X, W^*(X))$ also have a simple form:

$$\nabla_{x_i} F(X, W^*(X)) = \int_{GD(W^*)_i} \nabla_{x_i} d(x, x_i) \mathrm{d}\mu.$$
(9)

The pseudocode of our method is given in Algorithm 1. It is worth noting that for general cost kernel that is different from the 2-Wasserstein metric whose kernel is the squared distance, instead of using Newton's method, we use the L-BFGS method to update W, because the Hessian matrix is much more complicated in this case than that in the case.

- **2 Input:** domain D, density ρ , cost kernel d, number of points n, capacity constraints $\{c_i\}$ and a threshold ϵ as the termination condition.
- 3 Initialization: set k = 0 and X_0 to be *n* randomly generated sites. 4 repeat *//L-BFGS*
- **repeat** *//To meet the capacity constraints*
- 6 Update W_k by Newton's method or L-BFGS method
- 7 **until** $||\nabla_W F(X, W)|| < 10^{-12}$
- 8 Compute the gradients $\nabla_X F(X, W^*(X))$ in Eqn. (9)
- 9 Estimate the step-size for updating X by a line-search

```
10 X_{k+1} \leftarrow X_k
```

1

11 $k \leftarrow k+1$

12 until $||\nabla_X F(X, W^*(X))|| < \epsilon$

13 Output: (X_k, W_k)

4.2 Optimization Options

The computation of W^* is crucial to accurate computation of $\nabla_{x_i} F(X, W^*(X))$. As de Goes et al. [2012] observed, it usually takes 3 to 5 iterations to bring the residual of capacity constraints to be within an accuracy of 10^{-12} . Therefore we require $||\nabla_W F(X, W)|| < 10^{-12}$ during the step of updating W. In implementation, rather than always resetting W to 0 at the very beginning of Newton's method, it is better to use the previous values of W to initialize Newton's method. This is helpful for reducing the computation cost especially when $||\nabla_W F(X, W)||$ becomes very small. We observed an improvement of about 10% due to this "trick" in our experiments. When empty cells occur, we reduce the step size by half repeatedly until each site dominates a nonempty cell, as in [de Goes et al. 2012; Zhao et al. 2013]. Note that the convergence of this Newton step-control scheme was recently proved [Kitagawa et al. 2016].

For the 2-Wasserstein metric, the optimal partition is given by power diagrams. In this case, the Hessian matrix of F(X, W) w.r.t. Whas an elegant form [de Goes et al. 2012]. But for a general cost kernel different from the squared Euclidean distance, the coefficients of the Hessian matrix (gradient of the distance function integrated over the facets of the power diagram) are much more complicated, and in most cases not known in closed form. For this reason we use the L-BFGS method that only needs the first-order derivatives (that always correspond to the difference between the capacity constraints and the measures of the Laguerre cells).

4.3 Numerical Integration and Boundary Treatment

When the density function ρ is constant and the cost kernel is the 2-Wasserstein metric, the integrand in Eqn. (7) has an explicit representation by using the Green formula [Dobrovolsky 1972] to transform the computation of the cost function, as well as its gradients, into contour integration. But



in general cases, the cost function cannot be reduced to an explicit representation and thus discretization of the domain, typically a sufficiently refined triangulation, is required. Therefore, at the very beginning of the algorithm, we randomly distribute a set of points inside the domain and optimize their positions [Liu et al. 2009]. After that, we project the outer-ring points (whose Voronoi cells intersect the boundary) onto the boundary and re-optimize the other points. Finally we compute the constrained Delaunay triangulation w.r.t. these points as discretization of the domain; See the inset figure.

Besides, the optimal partition of the domain for a general cost kernel $d(\cdot, \cdot)$ may not be the power diagram. We have to develop an effective technique to build the χ_i partition with the help of the base



triangulation. First of all, we take the sites in X as sources and compute the weighted discrete distance field using the fast sweeping technique so that each vertex v of the base triangulation finds its nearest site x_i in X, i.e. $d(v, x_i) - w_i \le d(v, x_j) - w_j, \forall j \ne i$. After that, we compute the decomposition of each triangle depending on the following cases:

- \diamond If the site x_i gives the minimum weighted distances to v_1, v_2, v_3 at the same time, we simply assign the triangle f to x_i .
- ◇ If the site x_i gives the minimum weighted distance to v_1 and the site x_j gives the minimum weighted distances to v_2, v_3 , we come to find two dividing points $p_1 \in \overline{v_1 v_2}$ and $p_2 \in \overline{v_1 v_3}$ such that $\overline{p_1 p_2}$ can be taken as a segment delimiting x_i 's cell and x_j 's cell. The dividing point p_1 can be found from the equation $d(x_i, v_1) - w_i + (x - v_1) \cdot \nabla_x d(x, x_i) |_{v_1} = d(x_j, v_2) - w_j + (x - v_2) \cdot \nabla_x d(x, x_j) |_{v_2}$. The other dividing point p_2 can be computed likewise. See the inset figure.
- \diamond If the minimum weighted distances to v_1, v_2, v_3 are respectively given by x_i, x_j, x_k , then we find three dividing points $p_1 \in \overline{v_1 v_2}, p_2 \in \overline{v_2 v_3}, p_3 \in \overline{v_3 v_1}$ as mentioned above. We denote the center point by $p = (p_1 + p_2 + p_3)/3$. Finally, we assign the subcell $v_1 p_1 p p_3$ to $x_i, v_2 p_2 p p_1$ to x_j , and $v_3 p_3 p p_2$ to x_k , as the inset figure shows.

Another implementation detail we need to mention is boundary treatment. During the optimization process, some sites may jump out of the domain Ω and we need to pull them back to $\partial\Omega$ or the interior side of Ω . Suppose that in the *j*-th iteration, the *i*-th site $x_i^{(j)}$ is outside Ω and we wish to pull $x_i^{(j)}$ back. We find the intersection point *p* between $\partial\Omega$ and the connecting line segment of $x_i^{(j)}$ and its previous position $x_i^{(j-1)}$, and take *p* as the corrected position of $x_i^{(j)}$.

4.4 Performance Evaluation

Table 1: *Statistics of the number of operations of building power cells for the three examples shown in Fig. 2. Note that we take* $\|\nabla F\| \le 10^{-8}$ *as the termination condition.*

Examples	# of building power cells		Timing (s)	
	[de Goes et al. 2012]	Ours	[de Goes et al. 2012]	Ours
100 sites	4091	279	20.8	1.6
500 sites	2944	464	246.7	39.5
1000 sites	4307	471	556.0	58.4

We implemented and experimented with our CPD algorithm on a computer with a 64-bit version of Win7 system, a 3.07 GHz Intel(R) Core(TM) i7 CPU and 6 GB memory. The coding language is C++. Now we shall compare our CPD algorithm with the state of the art [de Goes et al. 2012]. For the examples shown in Fig. 2, we take $\|\nabla F\| \leq 10^{-8}$ as the termination condition. For the example of n = 100 sites (see the left column), we assume that the domain $[0, 1] \times [0, 1]$ is associated with a uniform density. For the



Figure 2: *Performance comparison with [de Goes et al., 2012]. The first row: snapshots of our algorithm. The second row: snapshots of [de Goes et al., 2012]. The third row: the plots of the cost function. The last row: the plots of the gradient of the cost function. Note that the horizontal axis in (g-l) denotes the timing (s).*

example of n = 500 sites (see the middle column), the density function at $x = (x^{(1)}, x^{(2)})$ is set to $\rho(x) = 0.1 + x^{(1)}$. For the example of n = 1,000 sites (see the right column), the density is set to $\rho(x) = e^{-8(x^{(1)}-0.5)^2-8(x^{(2)}-0.5)^2}$. Fig. 2(j-1) shows that our algorithm exhibits a faster convergence. In detail, we observe that (1) the resultant cells by our algorithm are of more compact shapes than [de Goes et al. 2012] after the same number of iterations; (2) the cost by our algorithm reaches the stable state earlier than [de Goes et al. 2012]; and (3) the gradient of the cost function decreases more sharply. Note that the horizontal axis of Fig. 2(g-1) denotes the timing.

Since the operation of building power cells is the most timeconsuming part, Table 2 shows the number of building power cells to achieve the same specified threshold 10^{-8} , as well as the total computation time. From the statistics we can see that our reduced gradient method has empirical super-linear convergence and significantly outperforms [de Goes et al. 2012].

4.5 General Cost Functions

Many real-life application occasions can be formulated as a CPD problem. Consider deploying some service centers (e.g. postal offices or medical centers) to serve the residents of a city, where each center has a capacity constraint. It's better to take the Euclidean distance, rather than the squared distance, as the service cost kernel. Another example is when the domain is a curved surface where we often take geodesic distances as the cost kernel. As pointed out in Section 3, our algorithmic framework supports general kernels. Here we use six typical kernels (see Fig. 3) to exhibit the extensibility of our algorithm, including

$$\diamond d(x, x_i) = (x^{(1)} - x_i^{(1)})^2 + (x^{(2)} - x_i^{(2)})^2 \text{ in } \mathbb{R}^2;$$

$$\diamond d(x, x_i) = \sqrt{(x^{(1)} - x_i^{(1)})^2 + (x^{(2)} - x_i^{(2)})^2} \text{ in } \mathbb{R}^2;$$

$$\diamond d(x, x_i) = (x^{(1)} - x_i^{(1)})^4 + (x^{(2)} - x_i^{(2)})^4 \text{ in } \mathbb{R}^2;$$

- ♦ Geodesic distance on mesh surfaces;
- ◊ Squared geodesic distance on mesh surfaces.
- ♦ Cubic geodesic distance on mesh surfaces.

In Fig. 3, the first row shows that CPD has freedom degrees to support user-specified capacity constraints while CVT cannot control cell sizes. The second row gives three CPDs with different cost kernels assuming the same uniform density. The third row shows the case of a Gaussian density function. The last row exhibits three CPDs on curved surfaces. When handling mesh surfaces, each site x_i is constrained to move along the tangent plane and projected onto the surface before evaluating the cost function. Furthermore, to find the optimal CPD partition, we have to frequently query the geodesic distances [Surazhsky and Surazhsky 2005; Xin and Wang 2009] between two points, which is computationally expensive. Due to this limitation, each of the input meshes in Fig. 3(i-k) contains about 1K triangle faces in our experiments. With the termination condi-tion $\|\nabla_W F(X)\|$ being less than 10^{-4} , we need about 15 iterations and 5 minutes to deal with each example. (Note that the partition of the surface is approximate and the evaluation of the cost function is approximate as well. That's why we set the tolerance to be 10^{-4} , rather than 10^{-8} .) However, this can be greatly accelerated using several recent methods for geodesic computation [Crane et al. 2013; Ying et al. 2013b; Ying et al. 2013a]. With a faster geodesic computation querier, we believe that it has a great potential for many geometry processing tasks.

5 Applications

In this section, we shall use it for three problems: displacement interpolation of function distributions, blue-noise point sampling, and optimal convex decomposition of 2D domains.

5.1 Displacement Interpolation

The traditional Wasserstein barycenter problem, i.e. displacement interpolation, provides a generic method for interpolating between distributions or functions based on advection instead of blending [Bonneel et al. 2011]. It has been found very useful in many applications such as shape interpolation [Solomon et al. 2015; Oberman and Ruan 2015] and texture synthesis [Bonneel et al. 2015]. The displacement interpolation is to find a target probability density ρ defined by

$$\min_{\rho} \sum_{k=1}^{m} \lambda_k \mathbf{W}_2^2(\rho_k, \rho), \tag{10}$$



(i) Geodesic distance (j) Squared (k) Cubic

Figure 3: Our algorithm can be extended to general cost kernels. The first row shows that CPD can generate compact cells with user-specified capacity constraints while CVT cannot. For the other examples, the capacity constraints divide the domain equally.

where $\{\rho_k\}_{k=1}^m$ are a collection of given probability densities and $\{\lambda_k\}_{k=1}^m$ are user-specified weights for computing the average probability density with regard to the 2-Wasserstein distance $\mathbf{W}(\cdot, \cdot)$. Recently many research works [Cuturi and Doucet 2013; Benamou et al. 2015; Solomon et al. 2015] suggested an energy term with an additional entropy regularization, bringing convexity to the problem and make it easier to solve. However, such an entropy term would result in an over-smoothed result in exchange for better efficiency. Besides, although lots of works proposed to compute the traditional Wasserstein barycenter distributions, few works aim at the semi-discrete setting. Here we propose a framework that supports multiple input distributions, each of which is represented by a collection of sites associated with capacities or a continuous probability density function. Our algorithm produces adaptively distributed samples to encode the final distribution.

Consider the following semi-discrete minimization problem, denoting $R_k = \{R_k^i\}_{i=1}^n : \Omega \to \{x_i\}_{i=1}^n$ as an arbitrary assignment between the support of ρ_k and X

$$\begin{cases} \min_{X} \sum_{k=1}^{m} \lambda_{k} \mathbf{W}_{2}^{2}(\rho_{k}, X, R_{k}), \\ s.t. \quad \int_{R_{k}^{i}} \rho_{k} \, \mathrm{d}x = c_{i}, \text{ for all } i, k. \end{cases}$$
(11)

The constraint term in (11) forces the mass transported to x_i under ρ_k to be equal to a given c_i . Based on the conclusion of [Aurenhammer et al. 1998], each assignment R_k minimizing the 2-Wasserstein cost under the capacity constraint should be a power diagram, and thus Eqn. (11) can be further reduced to the following minimization problem [de Goes et al. 2012]:

$$\begin{aligned} \text{Minimize } F_b(X) &= \sum_{k=1}^m \lambda_k \left(\sum_{i=1}^n \int_{GD(W)_k^i} \rho_k ||x - x_i||^2 \, \mathrm{d}x \\ &- \sum_{i=1}^n w_i (|GD(W)_k^i| - c_i) \right), \end{aligned}$$
(12)

with

$$\nabla_{x_i} F_b(x_i) = \sum_{k=1}^m 2c_i \lambda_k (x_i - b_k^i), \qquad (13)$$

where b_k^i is the centroid of the *i*-th power cell under the *k*-th distribution. In implementation, the optimization procedure can follow a similar procedure to computing CPDs, presented in Algorithm 2.

- 1 ____
- **2 Input:** densities $\{\rho_k\}_{k=1}^m$, weights $\{\lambda_k\}_{k=1}^m$, capacities $\{c_i\}_{i=1}^n$, and a threshold ϵ .
- **3 Output:** $\{x_i\}_{i=1}^n$
- 4 repeat *//L-BFGS*
- 5 for $k = 1, 2, 3, \cdots, m$
- 6 *//for the k-th distribution*
- 7 Optimize W_k until each power cell amounts to the given capacity
- 8 Compute the centroids of the power diagram under *k*-th distribution
- 9 end
- 10 Compute the cost function $F_b(X)$ in Eqn. (12)
- 11 Compute the gradient w.r.t. X; See Eqn. (13)
- 12 Update X by step search
- 13 until $||\nabla_X F_b(X)|| < \epsilon$

Algorithm 2: Capacity Constrained Wasserstein Barycenter with LBFGS.

Fig. 4 shows the result from two identical circular spots. The ideal distribution when the weights of these two inputs are both 0.5 should be of the same circular shape at the middle position. We compare our result with that by [Solomon et al. 2015] and visualize the color-coded errors in Fig. 4(c&g). The resolutions of both input images are 200×200 in this example. We can see that when n = 1.5K, we achieve an error of RMSE=0.078 in about 30 seconds, which is at the same efficiency/accuracy level with [Solomon et al. 2015]; But when n increases to 10K, the error of RMSE decreases to 0.0441 at the cost of more computation time - about 30 minutes for this example (the inefficiency is due to the density setting where lots of density values are very close to 0). Note that for comparison purpose, Fig. 4(e-f) are produced by an adaptive kernel estimation as the postprocessing [Abramson 1982]. Fig. 5 shows the results of 2-Wasserstein barycenters with multiple inputs.

5.2 Blue Noise Sampling

Stochastic point distributions with blue noise properties are widely used for various applications in computer graphics [Yan et al. 2015]. de Goes et al. [2012] presented a fast algorithm to generate highquality blue noise point distributions by finding a stationary point of the objective function defined in Eqn. (5). However, extremizing



Figure 4: Displacement interpolation of two identical circular spots. (a)&(d): the two inputs of circular spots: (b)&(c): interpo-

spots. (a)&(d): the two inputs of circular spots; (b)&(c): interpolated distribution by [Solomon et al. 2015] and the corresponding color-coded point-wise error (RMSE=0.0794), generated in 30 seconds (using the Matlab code provided by the authors of [Solomon et al. 2015]). (e-g) our results for n=1.5K and n=10K. When n = 1.5K, we achieve RMSE=0.078 in about 30 seconds; When n increases to 10K, the error of RMSE decreases to 0.0441 at the cost of increased computation time - about 30 minutes for this example.

the objective function does not prevent the regular patterns from emerging. In fact, the hexagonal grid patterns are the solutions to the extremization problem. de Goes et al. [2012] propose to jitter the points and run the optimization again when local regularities are detected. It is expected that a "shallower" extremum of the objective function which contains no regular patterns is obtained by a local searching algorithm. However, the optimization method of [de Goes et al. 2012] used a very loose stopping criteria, which usually terminates in 10 iterations, well before convergence. A more faithful local extremum can be efficiently achieved by our reduced gradient method that optimizes the same function. Experimental results show that "extreme" optimization of the objective function always results in structured point distributions even starting from a random point distribution (see Fig. 6(a)).

It is worth noting that all point samples are enforced to have the same capacity c in the existing methods based on capacityconstrained Voronoi tessellation [Balzer et al. 2009; Li et al. 2010; Chen et al. 2012; de Goes et al. 2012]. To prevent regular patterns in the point distributions, Zhang et al. [2016] suggested using non-uniform capacity constraints and dynamically adjust them during optimization. Inspired by this, we propose to jitter the capacity constraints for the points before optimization. A random noise with a maximum magnitude τ is used to jitter the capacity constraint for each point. The new capacity constraint c_i^* for each point is now specified by the following equation:

$$c_i = (1 + \tau \cdot rand(-1, 1))c, \quad c_i^* = \frac{c_i}{\sum c_i} nc,$$

where n is the sample number, and the second formula is to normalize the constraints so that they all sum to the total capacity of the domain.

Fig. 6(a) and 6(c) show point sets generated by extremizing the objective function with uniform capacity constraints and the jittered capacity constraints, respectively. A strict convergence threshold $(||\nabla_X F|| \le 10^{-7})$ is used for both cases. We can see that the jittered capacity constraints avoid regularity in the resultant point



Figure 6: Sampling results with different τ 's over a periodic domain with constant density. The blue noise properties can be analyzed through comparing the averaged periodograms, radial mean power (red curve), and anisotropy (blue curve).



Figure 5: The interpolation by our method of the four inputs $(500 \times 500 \text{ in size})$ at the four corners of the array (highlighted in red). Each output frame is generated by 10K sites in about 15 minutes.

distribution, producing high-quality blue noise. For the sake of fast computation, we set τ as a small number 0.2 and use a loose stopping condition with $\|\nabla_X F\| \le 0.05\sqrt{nm^3}$ in practice. Under this setting, our optimization method is able to converge after a dozen of iterations, and generate high-quality blue noise samplings, as shown in Fig. 6(c).

As we jitter the capacity constraints for sampling points, an immediate concern is that the point distribution may not adapt to the given density function precisely. Fig. 7 shows a sampling result generated by our method for an intensity ramp. We count the point density in each quarter of the ramp, and compare it to the reference density of the ramp. We can see that jittering the capacity constraints break local structured patterns while returns good adaption to the given density function.

Fig. 8 shows two image stippling results generated by our optimization framework. The sampling points $\{x_i\}_{i=1}^n$ are initialized randomly according to the piecewise-constant density function defined by the gray scale of the input image. Integrations involved in our formulation are exactly computed by constructing the pixel-cell intersections, as done in [de Goes et al. 2012].



Figure 7: Sampling of a quadratic density function with 1000 points, the percentages indicate point density in each quarter: (a) input quadratic density function; (b) sampling result by [de Goes et al. 2012]; (c) our result.

5.3 Centroidal Convex Decomposition

Another important problem in geometric computing, especially in mesh generation, is the convex decomposition of a geometric domain [Greene 1983; Ghosh et al. 2013], that is, decomposing a general non-convex domain into the union of a number of convex cells. The problem is relevant to numerous applications in pattern recognition, Minkowski sum computation, motion planning, computer graphics, and origami folding [Ghosh et al. 2013]. In this section we use the optimal capacity-constrained partition to provide a novel and effective solution to the convex decomposition of a 2D planar region.

Existing algorithms for convex decomposition usually minimize the number of convex components; See Fig. 9(a). In this paper, we propose a different type of convex decomposition, called the *optimal convex decomposition* that meets the following criteria: (1) each decomposed cell is convex; (2) the decomposed cells have specified areas; and (3) the decomposed cells are of compact shapes. Fig. 9(b) shows such a convex decomposition. Note that the term "optimal" is used to emphasize shape regularity rather than the number of decomposed cells.

Suppose that the boundary of the input 2D domain to be decom-



Figure 8: Image stippling. (a) &(c) Input images; (b) &(d) stippling results with 10K points generated by our optimization framework in 129 seconds and 146 seconds, respectively.



Figure 9: Convex decomposition of a star shape. The optimal convex decomposition (b) is much different than the traditional convex decomposition (a).

posed is defined by a polygonal curve. It is observed that simply running a CPD algorithm would cause a concave vertex on the boundary to be located in the interior of a single CPD cell, resulting in a non-convex cell (see Fig. 10(a)). Our key idea is to define all concave boundary vertices as *virtual sites* that are fixed and associated with a sufficiently small capacity constraint. Thus, the new cost function becomes

$$F(X,W) = \sum_{i=1}^{n+k} \int_{R_i^W} \rho(x) \|x - x_i\|^2 \mathrm{d}x$$
(14)

subject to

$$c_i = \int_{R_i^W} \rho(x) \mathrm{d}x (\triangleq m_i), 1 \le i \le n+k, \tag{15}$$

where n is the number of variable sites and k is the number of virtual sites on the boundary. Here $\{m_i, 1 \le i \le n\}$ is a set of capacity constraints for the moveable sites, and the number of such sites, n, is usually specified by the user. Let m_{average} be the average value of $\{m_i, 1 \le i \le n\}$. The capacity constraint for the virtual sites, say, m_{virtual} , is given by

$$m_{\text{virtual}} = \epsilon \times m_{\text{average}},$$
 (16)

where ϵ is set to be 10^{-3} in our experiments.

Furthermore, we assign a large multiplicative penalty to the cost terms defined by the virtual sites such that those small-sized power cells can be made compact and stay around the virtual sites (see Fig. 10(b)). The new cost function is as follows.

$$F(X,W) = \sum_{i=1}^{n+k} \int_{R_i^W} \rho(x) \|x - x_i\|^2 dx + \lambda \sum_{i=n+1}^{n+k} \int_{R_i^W} \rho(x) \|x - x_i\|^2 dx$$
(17)



Figure 10: Computation of the optimal convex decomposition: (a) Generally the concave points may be covered totally by a single CPD cell, giving rise to a non-convex cell. (b) Even if we enforce a small capacity for the virtual site at the concave point, the power cell stay non-convex after optimization. (c) After adding a large penalty coefficient λ for the virtual sites, the minimization makes all the normal power cells almost convex. (d) By removing the virtual sites from the results in (c), we obtain a convex decomposition of the domain.

Empirically we take the penalty coefficient to be

$$\lambda = \frac{m_{\text{normal}}}{m_{\text{average}}}.$$
(18)

After the above cost function is minimized (see Fig. 10(c)), we remove those virtual sites to get a clean convex decomposition in which each concave point is located on the common boundary of two adjacent convex power cells (see Fig. 10(d)). Fig. 11 demonstrates three examples of convex decomposition produced by our method based on CPDs with various density settings. We note that the density value should be larger in narrow regions and around concave points, generally implying a denser site distribution, to achieve a successful decomposition.

6 Conclusions and Future Works

In this paper, a super-linear convergent method is developed for computing the capacity-constrained centroidal power diagram. Experimental results show that our algorithm outperforms the existing method. Due to its effectiveness, we apply our fast CPD algorithm to three problems in computer graphics and geometric processing: displacement interpolation, blue-noise point sampling, and optimal



Figure 11: Convex decomposition of complex 2D shapes. The blue dots denote the user-specific concave points on the given boundary. For the Tortoise shape, we set a uniform density distribution. For the Guitar shape (b) and the Gear-Teeth shape (c), the density is higher in narrow regions or around the user-specified concave points.

convex decomposition of 2D domains. Furthermore, we show that the proposed algorithm can be extended to the capacity-constrained optimal partition with respect to general cost functions.

Our algorithm, in its current form, has several limitations. First, the capacity constraints are required to be met throughout the optimization process, which may be a cause of inefficiency. It is therefore valuable to develop an optimization technique that can accommodate an infeasible start. Second, the optimal mass transport problem in 3D space has been studied [Su et al. 2016; Lévy 2014]. We are therefore interested in extending our method to 3D or spaces of higher dimensions. Finally, we have only studied optimal convex decomposition of a 2D domain. The optimal convex decomposition of 3D domain is more relevant problem in practice and is much more challenging that the 2D version, so would need further study.

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7 Appendix

Theorem 1: For any fixed sites $X = \{x_i\}$, the capacityconstrained partition $R = \{R_i\}_{i=1}^n$ of the domain Ω that minimizes the cost function in Eqn. (4) is the one such that the boundary of any two adjacent regions R_a and R_b , which are associated with the sites a and b in X respectively, is define by the equation $d(x, a) - w_a = d(x, b) - w_b$ for some constants w_a and w_b .

PROOF: We shall prove the theorem by contradiction. Let us denote partition that minimizes the cost function in Eqn. (4). Assume that the theorem is not true. Then there exist two neighboring sites a and b with the shared boundary Γ of R_a and R_b such that there exist two distinct points $x_0, y_0 \in \Gamma$ for which

$$d(x_0, a) - d(x_0, b) \neq d(y_0, a) - d(y_0, b).$$

Without loss of generality, suppose that

$$d(x_0, a) - d(x_0, b) > d(y_0, a) - d(y_0, b)$$

Let $B(x, \delta)$ denote the ball centered at x with radius δ . Since d(x, y) is continuous with respect to x and y, there exist two balls $B(x_0, \delta_0)$ and $B(y_0, \delta_0)$ with sufficiently small $\delta_0 > 0$ such that

$$d(u,a) - d(u,b) > d(v,a) - d(v,b)$$

for all u in $B(x_0, \delta_0)$ and all v in $B(y_0, \delta_0)$.

Because the boundary Γ goes through the centers of $B(x_0, \delta_0)$ and $B(y_0, \delta_0)$, there exist two balls $B_x \equiv B(x_1, \delta_1)$ and $B_y \equiv B(y_1, \delta_2)$ with sufficiently small $\delta_1, \delta_2 > 0$ such that B_x and B_y have equal capacities (i.e. $\int_{u \in B_x} \rho(u) d\sigma = \int_{u \in B_y} \rho(v) d\sigma$) and that $B_x \subset R_a \cap B(x_0, \delta_0)$ and $B_y \subset R_b \cap B(y_0, \delta_0)$.

Since $B_x \subset B(x_0, \delta_0)$ and $B_y \subset B(y_0, \delta_0)$, we have

$$d(u, a) - d(u, b) > d(v, a) - d(v, b)$$

for all u in B_x and all v in B_y . It follows that

$$\int_{u\in B_x} \rho(u)(d(u,a) - d(u,b)) \mathrm{d}\sigma > \int_{v\in B_y} \rho(v)(d(v,a) - d(v,b)) \mathrm{d}\sigma,$$

which, after rearranging, yields

$$\int_{u \in B_x} \rho(u)d(u,a)d\sigma + \int_{v \in B_y} \rho(v)d(v,b)d\sigma$$

>
$$\int_{u \in B_x} \rho(u)d(u,b)d\sigma + \int_{v \in B_y} \rho(v)d(v,a)d\sigma \quad (19)$$

On the other hand, it follows from the optimality of R, in particular, the optimality of R_a and R_b that swapping B_x and B_y between R_a and R_b , while maintaining the capacities of R_a and R_b , does not reduce the total cost. Therefore

$$\int_{u \in B_x} \rho(u) d(u, a) \mathrm{d}\sigma + \int_{v \in B_y} \rho(v) d(v, b) \mathrm{d}\sigma$$
$$\leq \int_{u \in B_x} \rho(u) d(u, b) \mathrm{d}\sigma + \int_{v \in B_y} \rho(v) d(v, a) \mathrm{d}\sigma \quad (20)$$

Clearly, Eqn. 20 and Eqn. 19 contradict. Therefore we conclude that

$$d(x_0, a) - d(x_0, b) = d(y_0, a) - d(y_0, b),$$

for any two arbitrary points x_0 and y_0 on the boundary Γ . Hence, we have proved that the boundary curve Γ separating R_a and R_b is defined by an equation of the form

$$d(x,a) - w_a = d(x,b) - w_b$$
(21)

for some constants w_a and w_b . This completes the proof.

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