Abstract

We introduce a theoretical and algorithmic framework for multi-way graph partitioning that relies on a multiplicative cut-based objective. We refer to this objective as the Product Cut. We provide a detailed investigation of the mathematical properties of this objective and an effective algorithm for its optimization. The proposed model has strong mathematical underpinnings, and the corresponding algorithm achieves state-of-the-art performance on benchmark data sets.

1 Introduction

We propose the following model for multi-way graph partitioning. Let $G = (V, W)$ denote a weighted graph, with $V$ its vertex set and $W$ its weighted adjacency matrix. We define the Product Cut of a partition $P = (A_1, \ldots, A_R)$ of the vertex set $V$ as

$$
Pcut(P) = \prod_{r=1}^{R} \frac{Z(A_r, A_r^c)}{e^{H(P)}}, \quad H(P) = -\sum_{r=1}^{R} \theta_r \log \theta_r,
$$

where $\theta_r = |A_r|/|V|$ denotes the relative size of a set. This model provides a distinctive way to incorporate classical notions of a quality partition. The non-linear, non-local function $Z(A_r, A_r^c)$ of a set measures its intra- and inter-connectivity with respect to the graph. The entropic balance $H(P)$ measures deviations of the partition $P$ from a collection of sets $(A_1, \ldots, A_R)$ with equal size. In this way, the Product Cut optimization parallels the classical Normalized Cut optimization in terms of its underlying notion of cluster, and it arises quite naturally as a multiplicative version of the Normalized Cut.

Nevertheless, the two models strongly diverge beyond the point of this superficial similarity. We provide a detailed analysis to show that (1) settles the compromise between cut and balance in a fundamentally different manner than classical objectives, such as the Normalized Cut or the Cheeger Cut. The sharp inequalities

$$
0 \leq \text{Ncut}(P) \leq 1 \quad e^{-H(P)} \leq \text{Pcut}(P) \leq 1
$$

succinctly capture this distinction; the Product Cut exhibits a non-vanishing lower bound while the Normalized Cut does not. We show analytically and experimentally that this distinction leads to superior stability properties and performance. From an algorithmic point-of-view, we show how to cast the minimization of (1) as a convex maximization program. This leads to a simple, exact continuous relaxation of the discrete problem that has a clear mathematical structure. We leverage this formulation to develop a monotonic algorithm for optimizing (1) via a sequence of linear programs, and we introduce a randomized version of this strategy that leads to a simple yet highly effective...
We begin by introducing our notation and by describing the rationale underlying our model. We use $G = (V, W)$ to denote a graph on $n$ vertices $V = \{v_1, \ldots, v_n\}$ with weighted edges $W = \{w_{ij}\}_{i,j=1}^n$ that encode similarity between vertices. We denote partitions of the vertex set into $R$ subsets as $\mathcal{P} = (A_1, \ldots, A_R)$, with the understanding that the $A_r \subset V$ satisfy the covering $A_1 \cup \cdots \cup A_R = V$ constraint, the non-overlapping $A_r \cap A_s = \emptyset$, $(r \neq s)$ constraint and the non-triviality $A_r \neq \emptyset$ constraint. We use $f, g, h, u, v$ to denote vertex functions $f : V \to \mathbb{R}$, which we view as functions $f(v_i)$ and $n$-vectors $f \in \mathbb{R}^n$ interchangeably. For a $A \subset V$ we use $|A|$ for its cardinality and $I_A$ for its indicator function. Finally, for a given graph $G = (V, W)$ we use $D := \text{diag}(W1_V)$ to denote the diagonal matrix of weighted vertex degrees.

The starting point for our model arises from a well-known and widely used property of the random walk on a graph. Namely, a random walker initially located in a cluster $A$ is unlikely to leave that cluster quickly [8]. Different approaches of quantifying this intuition then lead to a variety of multi-way partitioning strategies for graphs [11, 12, 1]. The personalized page-rank methodology provides an example of this approach. Following [1], given a scalar $0 < \alpha < 1$ and a non-empty vertex subset $A$ we define

$$\text{pr}_A := M_\alpha^{-1}1_A/|A| \quad M_\alpha := (I - \alpha WD^{-1}) / (1 - \alpha)$$

as its personalized page-rank vector. As $1_A/|A|$ is the uniform distribution on the set $A$ and $WD^{-1}$ is the transition matrix of the random walk on the graph, $\text{pr}_A$ corresponds to the stationary distribution of a random walker that, at each step, moves with probability $\alpha$ to a neighboring vertex by a usual random walk, and has a probability $(1 - \alpha)$ to teleport to the set $A$. If $A$ has a reasonable cluster structure, then $\text{pr}_A$ will concentrate on $A$ and assign low probabilities to its complement. Given a high-quality partition $\mathcal{P} = (A_1, \ldots, A_R)$ of $V$, we therefore expect that $\sigma_A := \text{pr}_{A_r}(v_i)$ should achieve its maximal value over $1 \leq r \leq R$ when $r = r(i)$ is the class of the $i$th vertex.

Viewed from this perspective, we can formulate an $R$-way graph partitioning problem as the task of selecting $\mathcal{P} = (A_1, \ldots, A_R)$ to maximize some combination of the collection $\{\sigma_i, r(i) : i \in V\}$ of page-rank probabilities generated by the partition. Two intuitive options immediately come to mind, the arithmetic and geometric means of the collection:

$$\text{Maximize} \quad \frac{1}{n} \sum_r \sum_{v_i \in A_r} \text{pr}_{A_r}(v_i) \quad \text{over all partitions } (A_1, \ldots, A_R) \text{ of } V \text{ into } R \text{ sets.}$$

$$\text{Maximize} \quad \left(\prod_r \prod_{v_i \in A_r} \text{pr}_{A_r}(v_i)^{1/n}\right)^{1/n} \quad \text{over all partitions } (A_1, \ldots, A_R) \text{ of } V \text{ into } R \text{ sets.}$$

The first option corresponds to a straightforward variant of the classical Normalized Cut. The second option leads to a different type of cut-based objective that we term the Product Cut. The underlying reason for considering [5] is quite natural. If we view each $\text{pr}_{A_r}$ as a probability distribution, then [5] corresponds to a formal likelihood of the partition. This proves quite analogous to re-formulating the classical $k$-means objective for partitioning $n$ data points $(x_1, \ldots, x_n)$ into $R$ clusters $(A_1, \ldots, A_R)$ in terms of maximizing a likelihood

$$\prod_{r=1}^R \prod_{v_i \in A_r} \exp\left(-\frac{||x_i - m_r||^2}{2\sigma_r^2}\right)$$

of Gaussian densities. While the Normalized Cut variant [4] is certainly popular, we show that it suffers from several defects that the Product Cut resolves. As the Product Cut can be effectively optimized and generally leads to higher quality partitions, it therefore provides a natural alternative.

To make these ideas precise, let us define the $\alpha$-smoothed similarity matrix as $\Omega_\alpha := M_\alpha^{-1}$ and use $\{\omega_{ij}\}_{i,j=1}^n$ to denote its entries. Thus $\omega_{ij} = (M_\alpha^{-1}1_{v_i})_i = \text{pr}_{A_r}(v_i)$, and so $\omega_{ij}$ gives a non-local measure of similarity between the vertices $v_i$ and $v_j$ by means of the personalized page-rank diffusion process. The matrix $\Omega_\alpha$ is column stochastic, non-symmetric, non-sparse, and has diagonal entries.
Moreover the lower bound is attained for partitions which choose the partition that maximizes the entropic balance. So in this case, any partition The Normalized Cut does not possess this property. As an extreme but easy-to-analyze example, The lower bound in (9) can be directly read from (6) and (7), while the upper bound is non-trivial and as its Product Cut and Normalized Cut, respectively. The non-linear, non-local function properties using experimental data as well as a simplified model problem that isolates the source of perturbations. We now delve deeper and contrast the two objectives by analyzing their stability properties.

2.1 (In-)Stability Properies of the Product Cut and Normalized Cut

In practice, the stronger balancing effect of the Product Cut manifests as a stronger tolerance to perturbations. We now delve deeper and contrast the two objectives by analyzing their stability properties using experimental data as well as a simplified model problem that isolates the source of
The inherent difficulties. Invoking ideas from dynamical systems theory, we say an objective is \textit{stable} if an infinitesimal perturbation of a graph \( G = (V, W) \) leads to an infinitesimal perturbation of the optimal partition. If an infinitesimal perturbation leads to a dramatic change in the optimal partition, then the objective is \textit{unstable}.

We use a simplified model to study stability of the Product Cut and Normalized Cut objectives. Consider a graph \( G_n = (V_n, W_n) \) made of two clusters \( A_n \) and \( B_n \) containing \( n \) vertices each. Each vertex in \( G_n \) has degree \( k \) and is connected to \( k \) vertices in the opposite cluster, where \( 0 \leq \mu \leq 1 \).

The graph \( G_n^{0} \) is a perturbation of \( G_n \), constructed by adding a small cluster \( C \) of size \( n_0 \ll n \) to the original graph. Each vertex of \( C \) has degree \( k_0 \) and is connected to \( \mu k_0 \) vertices in \( B_n \) and \( (1 - \mu k_0) k_0 \) vertices in \( C \) for some \( 0 \leq \mu_0 \leq 1 \). In the perturbed graph \( G_n^{0} \), a total of \( n_0 \) vertices in \( B_n \) are linked to \( C \) and have degree \( k + \mu_0 k_0 \).

The supplementary material contains the proof.

We summarize the conclusions of this analysis in the theorem below, which shows that the Normalized Cut is unstable in certain parameter regimes while the Product Cut is always stable. The supplementary material contains the proof.

**Theorem 2** Suppose that \( \mu, \mu_0, k, k_0, n_0 \) are fixed. Then

\[
\mu_0 < 2\mu \quad \Rightarrow \quad \text{Ncut}_{G_n^{0}}(P^{0, \text{good}}_n) > \text{Ncut}_{G_n^{0}}(P^{0, \text{bad}}_n) \quad \text{for } n \text{ large enough}. \quad (11)
\]

\[
\text{Pcut}_{G_n^{0}}(P^{0, \text{good}}_n) < \text{Pcut}_{G_n^{0}}(P^{0, \text{bad}}_n) \quad \text{for } n \text{ large enough}. \quad (12)
\]

Statement (11) simply says that the large cluster \( A_n \) must have a conductance \( \mu \) at least twice better than the conductance \( \mu_0 \) of the small perturbation cluster \( C \) in order to prevent instability. Thus adding an infinitesimally small cluster with mediocre conductance (up to two times worse the conductance of the main structure) has the potential of radically changing the partition selected by the Normalized Cut. Moreover, this result holds for the classical Normalized Cut, its smoothed variant (4) as well as for similar objectives such as the Cheeger Cut and Ratio Cut. Conversely, (12) shows that adding an infinitesimally small cluster will not affect the partition selected by the
Our strategy for optimizing the Product Cut relies on a popular paradigm for discrete optimization, i.e., exact relaxation. We begin by showing that the discrete, graph-based formulation (5) can be relaxed to a continuous optimization problem, specifically a convex maximization program. We then prove that this relaxation is exact, in the sense that optimal solutions of the discrete and continuous problems coincide. With an exact relaxation in hand, we may then appeal to continuous optimization strategies (rather than discrete or greedy ones) for optimizing the Product Cut. This general idea of exact relaxation is intimately coupled with convex maximization.

Assume that the graph \( G = (V, W) \) is connected. Then by taking the logarithm of (5) we see that (5) is equivalent to the problem

\[
\text{Maximize } \sum_{r=1}^{R} \sum_{A_r} \log \left( \frac{\Omega_n 1_{A_r}}{|A_r|} \right)
\]

over all partitions \( \mathcal{P} = (A_1, \ldots, A_R) \) of \( V \) into \( R \) non-empty subsets. 

\[
(\text{P})
\]
The relaxation of (P) then follows from the usual approach. We first encode sets \( A_r \subseteq V \) as binary vertex functions \( 1_{A_r} \), then relax the binary constraint to arrive at a continuous program. Given a vertex function \( f \in \mathbb{R}_+^n \) with non-negative entries, we define the continuous energy \( e(f) \) as

\[
e(f) := \langle f, \log \left( \Omega f / \langle f, 1_V \rangle \right) \rangle \quad \text{if } f \neq 0, \quad \text{and } \ e(0) = 0,
\]

where \( \langle \cdot, \cdot \rangle \) denotes the usual dot product in \( \mathbb{R}^n \) and the logarithm applies entrywise. As \( (\Omega f)_i > 0 \) whenever \( f \neq 0 \), the continuous energy is well-defined. After noting that \( \sum_r e(1_{A_r}) \) is simply the objective value in problem \( (P) \), we arrive to the following continuous relaxation

\[
\begin{align*}
\text{Maximize } & \sum_{r=1}^R e(f_r) \\
\text{over all } & (f_1, \ldots, f_R) \in \mathbb{R}_+^n \times \ldots \times \mathbb{R}_+^n \text{ satisfying } \sum_{r=1}^R f_r = 1_V
\end{align*}
\]

(P-rlx)

where the non-negative cone \( \mathbb{R}_+^n \) consists of all vectors in \( \mathbb{R}^n \) with non-negative entries.

The following theorem provides the theoretical underpinning for our algorithmic approach. It establishes convexity of the relaxed objective for connected graphs.

**Theorem 3** Assume that \( G = (V, W) \) is connected. Then the energy \( e(f) \) is continuous, positive 1-homogeneous and convex on \( \mathbb{R}_+^n \). Moreover, the strict convexity property

\[
e(\theta f + (1 - \theta) g) < \theta e(f) + (1 - \theta) e(g) \quad \text{for all } \theta \in (0, 1)
\]

holds whenever \( f, g \in \mathbb{R}_+^n \) are linearly independent.

The continuity of \( e(f) \) away from the origin as well as the positive one-homogeneity are obvious, while the continuity of \( e(f) \) at the origin is easy to prove. The proof of convexity of \( e(f) \), provided in the supplementary material, is non-trivial and heavily relies on the particular structure of \( \Omega f \) itself. With convexity of \( e(f) \) in hand, we may prove the main theorem of this section.

**Theorem 4** (Equivalence of (P) and (P-rlx)) Assume that \( G = (V, W) \) is connected and that \( V \) contains at least \( R \) vertices. If \( P = (A_1, \ldots, A_R) \) is a global optimum of \( (P) \) then \( (1_{A_1}, \ldots, 1_{A_R}) \) is a global optimum of \( (P-rlx) \). Conversely, if \( (f_1, \ldots, f_R) \) is a global optimum of \( (P-rlx) \) then \( (f_1, \ldots, f_R) = (1_{A_1}, \ldots, 1_{A_R}) \) where \( (A_1, \ldots, A_R) \) is a global optimum of \( (P) \).

**Proof.** By strict convexity, the solution of the maximization (P-rlx) occurs at the extreme points of the constraint set \( \Sigma = \{(f_1, \ldots, f_R) : f_r \in \mathbb{R}_+^n \text{ and } \sum_{r=1}^R f_r = 1\} \). Any such extreme point takes the form \( (1_{A_1}, \ldots, 1_{A_R}) \), where necessarily \( A_1 \cup \ldots \cup A_R = V \) and \( A_r \cap A_s = \emptyset (r \neq s) \) hold. It therefore suffices to rule out extreme points that have an empty set of vertices. But if \( A \neq B \) are non-empty then \( 1_A, 1_B \) are linearly independent, and so the inequality \( e(1_A + 1_B) < e(1_A) + e(1_B) \) holds by strict convexity and one-homogeneity. Thus given a partition of the vertices into \( R - 1 \) non-empty subsets and one empty subset, we can obtain a better energy by splitting one of the non-empty vertex subsets into two non-empty subsets. Thus any globally maximal partition cannot contain empty subsets. \( \square \)

With theorems 3 and 4 in hand, we may now proceed to optimize \( (P) \) by searching for optima of its exact relaxation. We tackle the latter problem by leveraging sequential linear programming or gradient thresholding strategies for convex maximization. We may write (P-rlx) as

\[
\begin{align*}
\text{Maximize } & \mathcal{E}(F) \quad \text{subject to } \ F \in C \quad \text{and } \ \psi_i(F) = 0 \text{ for } i = 1, \ldots, n \quad (13)
\end{align*}
\]

where \( F = (f_1, \ldots, f_R) \) is the optimization variable, \( \mathcal{E}(F) \) is the convex energy to be maximized, \( C \) is the bounded convex set \([0, 1]^n \times \ldots \times [0, 1]^n \) and the \( n \) affine constraints \( \psi_i(F) = 0 \) correspond to the row-stochastic constraints \( \sum_{r=1}^R f_{i,r} = 1 \). Given a current feasible estimate \( F^k \) of the solution, we obtain the next estimate \( F^{k+1} \) by solving the linear program

\[
\begin{align*}
\text{Maximize } & L_k(F) \quad \text{subject to } \ F \in C \quad \text{and } \ \psi_i(F) = 0 \text{ for } i = 1, \ldots, n \quad (14)
\end{align*}
\]

where \( L_k(F) = \mathcal{E}(F^k) + \langle \nabla \mathcal{E}(F^k), F - F^k \rangle \) is the linearization of the energy \( \mathcal{E}(F) \) around the current iterate. By convexity of \( \mathcal{E}(F) \), this strategy monotonically increases \( \mathcal{E}(F^k) \) since \( \mathcal{E}(F^{k+1}) \geq L_k(F^{k+1}) \geq L_k(F^k) = \mathcal{E}(F^k) \). The iterates \( F^k \) therefore encode a sequence of partitions of \( V \) that monotonically increase the energy at each step. Either the current iterate maximizes the linear form, in which case first-order optimality holds, or else the subsequent iterate produces a partition with a...
We conclude our study of the Product Cut model by presenting extensive experimental evaluation. When initialized from a random partition, the iterates almost immediately converge to a poor-quality solution. We may rescue this poor quality algorithm and convert it to a highly effective one, while maintaining its simplicity, by randomizing the LP (14) at each step in the following way. At step $k$ we solve the LP

$$\text{maximize } L_k(F) \quad \text{subject to } F \in C \text{ and } \psi_i(F) = 0 \text{ for } i \in \mathcal{I}_k,$$

where the set $\mathcal{I}_k$ is a random subset of $\{1, 2, \ldots, n\}$ obtained by drawing $s_k$ constraints uniformly at random without replacement. The LP (15) is therefore version of LP (14) in which we have dropped a random set of constraints. If we start by enforcing a small number $s_k$ of constraints and slowly increment this number $s_{k+1} = s_k + \Delta s_k$ as the algorithm progresses, we allow the algorithm time to explore the energy landscape. Enforcing more constraints as the iterates progress ensures that (15) eventually coincides with (14), so convergence of the iterates $F^k$ of the randomized algorithm is still guaranteed. The attraction is that LP (15) has a simple, closed-form solution given by a variant of gradient thresholding. We derive the closed form solution of LP (15) in section 1 of the supplementary material, and this leads to Algorithm 1 above.

The overall effectiveness of this strategy relies on two key ingredients. The first is a proper choice of the number of constraints $s_k$ to enforce at each step. Selecting the rate at which $s_k$ increases is similar, in principle, to selecting a learning rate schedule for a stochastic gradient descent algorithm. If $s_k$ increases too quickly then the algorithm will converge to poor-quality partitions. If $s_k$ increases too slowly, the algorithm will find a quality solution but waste computational effort. A good rule of thumb is to linearly increase $s_k$ at some constant rate $\Delta s_k \equiv \lambda$ until all constraints are enforced, at which point we switch to the deterministic algorithm and terminate the process at convergence. The second key ingredient involves approximating solutions to the linear system $M_s x = b$ quickly. We use a simple Algebraic Multigrid (AMG) technique, i.e. a stripped-down version of [7] or [6], to accomplish this. The main insight here is that exact solutions of $M_s x = b$ are not needed, but not all approximate solutions are effective. We need an approximate solution $x$ that has non-zero entries on all of $|V|$ for thresholding to succeed, and this can be accomplished by AMG at very little cost.

4 Experiments

We conclude our study of the Product Cut model by presenting extensive experimental evaluation of the algorithm$^1$. We intend these experiments to highlight the fact that, in addition to a strong theoretical model, the algorithm itself leads to state-of-the-art performance in terms of cluster purity on a variety of real world data sets. We provide experimental results on four text data sets (20NEWS, RCV1, WEBKB4, CITESEER) and four data sets containing images of handwritten digits (MNIST, PENDIGITS, USPS, OPTDIGITS). We provide the source for these data sets and details on their

$^1$The code is available at https://github.com/xbresson/pcut
Table 1: Algorithmic Comparison via Cluster Purity.

<table>
<thead>
<tr>
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<th>20NE</th>
<th>RCV1</th>
<th>WEBK</th>
<th>CITE</th>
<th>MNIS</th>
<th>PEND</th>
<th>USPS</th>
<th>OPTI</th>
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<td>3.3K</td>
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<td>43</td>
<td>96</td>
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<td>LSD</td>
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<td>53</td>
<td>76</td>
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<tr>
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<td>54</td>
<td>97</td>
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<td>87</td>
<td></td>
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<tr>
<td>PCut (.9, λ)</td>
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<td>58</td>
<td>63</td>
<td>97</td>
<td>87</td>
<td>89</td>
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<tr>
<td>PCut (.9, λ)</td>
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<td>50</td>
<td>57</td>
<td>64</td>
<td>96</td>
<td>84</td>
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<td>95</td>
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</table>

We compare our method against partitioning algorithms that, like the Product Cut, rely on graph-cut objective principles and that partition the graph in a direct, non-recursive manner. The NCut algorithm [15] is a widely used spectral algorithm that relies on a post-processing of the eigenvectors of the graph Laplacian to optimize the Normalized Cut energy. The NMFR algorithm [14] uses a graph-based random walk variant of the Normalized Cut. The LSD algorithm [2] provides a non-negative matrix factorization algorithm that relies upon a trace-based relaxation of the Normalized Cut objective. The MTV algorithm from [3] and the balanced k-cut algorithm from [9] provide total-variation based algorithms that attempt to find an optimal multi-way Cheeger cut of the graph by using ℓ1 optimization techniques. Both algorithms optimize the same objective and achieve similar purity values. We report results for [3] only. The GRACLUS algorithm [4, 5] uses a multi-level coarsening approach to optimize the NCut objective as formulated in terms of kernel k-means. Table 1 reports the accuracy obtained by these algorithms for each data set. We use cluster purity to quantify the quality of the calculated partition, defined according to the relation:

\[ \text{Purity} = \frac{1}{n} \sum_{r=1}^{R} \max_{1 \leq i < R} m_{r,i}. \]

Here \( m_{r,i} \) denotes the number of data points in the \( r \)-th cluster that belong to the \( i \)-th ground-truth class. The third row of the table (RND) provides a base-line purity for reference, i.e. the purity obtained by assigning each data point to a class from 1 to \( R \) uniformly at random. The PCut, MTV and GRACLUS algorithms rely on randomization, so for these algorithms we report the average purity achieved over 500 different runs. For the PCut algorithm, we use \( \alpha = .9 \) when defining \( \Omega_\alpha \). Also, in order to illustrate the tradeoff when selecting the rate at which the number of enforced constraints \( s_k \) increases, we report accuracy results for the linear rates

\[ \Delta s_k = 10^{-4} \times n := \lambda_1 \quad \text{and} \quad \Delta s_k = 5 \times 10^{-4} \times n := \lambda_2 \]

where \( n \) denotes the total number of vertices in the data set. By and large both PCut and NMFR consistently outperform the other algorithms in terms of accuracy.

Table 2: Computational Time

<table>
<thead>
<tr>
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<th>MNIST</th>
<th>20NEWS</th>
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<tbody>
<tr>
<td></td>
<td>NMFR</td>
<td>PCut (.9, λ1)</td>
</tr>
<tr>
<td></td>
<td>4.6mn</td>
<td>11s (92%)</td>
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<td></td>
<td>(92%)</td>
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In addition to the accuracy comparisons, table 2 records the time required for PCut and NMFR to reach 95% of their limiting purity value on the two largest data sets, 20NEWS and MNIST. Each algorithm is implemented in a fair and consistent way, and the experiments were all performed on the same architecture. Timing results on the smaller data sets from table 1 are consistent with those obtained for 20NEWS and MNIST. In general we observe that PCut runs significantly faster. Additionally, as we expect for PCut, the slower rate \( \lambda_1 \) generally leads to more accurate results while the larger rate \( \lambda_2 \) typically converges more quickly.

When taken together, our theoretical and experimental results clearly reveal that the model provides a promising method for graph partitioning. The algorithm consistently achieves state-of-the-art results, and typically runs significantly faster than other algorithms that achieve a comparable level of accuracy. Additionally, both the model and algorithmic approach rely upon solid mathematical foundations that are frequently missing in the multi-way clustering literature.

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References


