A Uniform Approach to Accelerated PageRank Computation

Frank McSherry
Microsoft Research, SVC
1065 La Avenida
Mountain View, CA, USA 94043
mcsherry@microsoft.com

ABSTRACT
In this note we consider a simple reformulation of the traditional power iteration algorithm for computing the stationary distribution of a Markov chain. Rather than communicate their current probability values to their neighbors at each step, nodes instead communicate only changes in probability value. This reformulation enables a large degree of flexibility in the manner in which nodes update their values, leading to an array of optimizations and features, including faster convergence, efficient incremental updating, and a robust distributed implementation.

While the spirit of many of these optimizations appear in previous literature, we observe several cases where this unification simplifies previous work, removing technical complications and extending their range of applicability. We implement and measure the performance of several optimizations on a sizable (34M node) web subgraph, seeing significant composite performance gains, especially for the case of incremental recomputation after changes to the web graph.

Categories and Subject Descriptors
H.3.3 [Information Storage and Retrieval]: Information Search and Retrieval

General Terms
Algorithms, Experimentation, Performance, Theory

Keywords
PageRank, web graph, link analysis, random walks

1. INTRODUCTION
Motivated largely by the success and scale of Google’s PageRank ranking function, much research has emerged on efficiently computing the stationary distributions of web-scale Markov chains, the mathematical mechanism underlying PageRank. The main challenge is that the web graph is so large that its edges typically only exist in external memory and an explicit representation of its stationary distribution just barely fits in to main memory. The time required to compute the stationary distribution is on the order of tens of hours to days, and constant factor improvements in running times can save substantial time and money. Even for the common researcher with interests in ranking research, computing and recomputing vectors of ranks is a time consuming processes that greatly limits research throughput.

As such, much work has been done on accelerating the performance of PowerIteration, the traditional approach to computing stationary distributions. These optimizations cover a spectrum of techniques, ranging from transformations to the Markov chain that accelerate mixing to efficient heuristic updates that behave like PowerIteration to clever reuse of previously computed solutions. Most of these techniques are developed and evaluated in isolation, and it is unclear to what degree they can be effectively combined, both in terms of implementation and performance.

1.1 Notation and Terminology
Throughout this note we will frequently refer to vectors, matrices, and the scalars they comprise. For clarity, we consistently use lowercase letters (x) for vectors and capital letters (A) for matrices. For each, subscripted quantities (x_u and A_uv) are used to reference the scalar values at the indexed coordinates.

1.2 PageRank and PowerIteration
PageRank [2] is a system of scoring nodes in a directed graph based on the stationary distribution of a random walk on the directed graph. Conceptually, the score of a node corresponds to the frequency with which the node is visited as an individual strolls randomly through the graph. For technical reasons, the random walk is also encouraged to occasionally reset to a prespecified distribution, overcoming issues of weakly connected components in which a random surfer might get stuck and accelerating the rate at which a random walk approaches the stationary distribution.

A random walk on n nodes can be described by a n × n matrix P, where entry P_{uv} is the probability that from node u the walk next arrives at node v. Starting from a distribution x over the nodes (x is a vector of n entries that are non-negative and sum to one), after one step the distribution becomes Px, and more generally after t steps the distribution becomes P^t x.

We can decompose P into those transitions due to traversing a web link, and those transitions due to random resetting. Let the sparse matrix A have entries A_{uv} equal to the probability that from node u the walk traverses the link (u, v) to node v. Additionally, we define the vector r with each coor-
dinate \( r_u = 1 - \sum_v A_{uv} \), equal to the probability that the
that the walk chooses not to follow an arc from node \( u \) and
instead resets randomly to a node \( v \) chosen with probability
proportional to \( d_v \). \( P \) can then be written as \( P = A + dr^T \),
capturing both of the types of transitions.

PowerIteration is the traditional manner of computing the
stationary distribution of \( P \), explicitly simulating the dis-
semination of probability mass by repeatedly applying
\( P \) to a supplied initial distribution \( x \). Under modest assump-
tions, e.g. that all entries of \( d \) and \( r \) are positive, for any
initial distribution \( x \), \( P^T x \) converges to a unique stationary
distribution as \( i \) increases.

**PowerIteration** \((P, x)\)

1. While (not converged)
   
   (a) Set \( x = Px \)

   While \( P \) itself is a dense matrix, every node can reset to
any other node, we can efficiently compute \( Px \) by viewing it as
\( Ax + dr^T x \). We assume that \( A \) is stored on disk in a sparse
format, perhaps as a list of \((source, target, value)\) triples,
though there are more compact representations. \( Ax \) is then
computed using sparse matrix-vector multiplication: since
\((Ax)_v = \sum_u A_{uv}x_u \), we can populate the result vector by
scanning the edge file, for each non-zero \( A_{uv} \) adding \( A_{uv}x_u \)
to coordinate \( v \) of the result. We can produce the vector
dr^T x \) by determining \( r^T x \) in a pass over \( r \) and \( x \), and scaling
d \) appropriately before adding it to \( Ax \).

For acceptable performance, we may only perform sequen-
tial access to the edge file, which is too large to fit into main
memory. Generally speaking, the number of non-zero entries
in \( A \) is the limiting performance factor, both because of our
need to scan over the edge file to read these entries, and also
the random accesses to \( x \) each entry requires. The other op-
erations, vector addition, scaling, and inner product, can all
be done using sequential access to main memory.

2. AN UPDATE-BASED ALGORITHM

Oddly, we start our generalization of PowerIteration by
restricting the problem we address. There are many vectors
satisfying \( x = Px \); any solution \( x \) can be multiplied by an
arbitrary scalar value and still satisfy the equality. Typically
we focus our attention on finding the vector \( x \) with \( \|x\|_1 = 1 \).
Let us instead focus on finding the vector \( x \) for which
\( r^T x = 1 \), and for which

\[
x = Px = Ax + dr^T x = Ax + d.
\]

As we will see in Theorem 1, if \( x = Ax + d \), then \( x = Px \).
Normalized, this vector is the stationary distribution of \( P \).

Consider an analog of PowerIteration in which we repeat-
eddy set \( x = Ax + d \). As with PowerIteration, this iterative
process will converge, and it converges to a vector satisfying
\( x = Ax + d \). We can monitor convergence through the
vector \( y = Ax - x + d \); so long as \( y \) is non-zero, \( x \) has not
yet converged. But, \( y \) also tells us the direction to update \( x \);
we advance to the next iterate of \( x \) by adding \( y \), yielding
\( Az + d \). This first role is crucial, we must bring \( y \) to zero, but
we needn’t be so rigid as to only ever add \( y \) to \( x \). We might
instead add other vectors to \( x \) that yield forward progress,
maintaining \( y \) both as a convergence criteria, but also for
guidance in choosing updates to \( x \).

Consider an algorithm that monitors \( y = Ax - x + d \), but
is free to choose an arbitrary update vector \( z \) at any step,
advancing from \( x \) to \( x + z \). It is not hard to appropriately
update \( y \), as its new value satisfies

\[
A(x + z) - (x + z) + d = y + Az - z.
\]

For any update vector \( z \), we can update \( y \) by passing \( z 
through the matrix \( A \), adding the result to \( y \), and subtract-
ing \( z \). Intuitively, \( z \) is extracted from \( y \) and propagated
across the links in \( A \), informing nodes of changes in their
parent’s values and insisting that they now update in turn.

Operationally, this is exactly the algorithmic framework
that we will consider. However, it will be useful not to fix
\( y \) in terms of a particular \( A, x, \) and \( d \), but rather let it be
specified as an input parameter, properly determined before
the method is invoked.

**UpdateIteration** \((A, x, y)\)

1. While (updates \( y \) remain):
   
   (a) Choose an update vector \( z \).
   (b) Set \( x = x + z \).
   (c) Set \( y = y + (Az - z) \).

While this framework is presently little more than a system
of bookkeeping, we will solidify how one might choose \( z \)
to shrink \( \|y\|_1 \), and which choices lead to efficient algorithms.

We now state two theorems regarding the limit and rate of
convergence of UpdateIteration \((A, x, y)\). The proofs, while
short, are rote and unilluminating, and are deferred to Ap-
pendix A. We first argue that choosing \( y = Ax - x + d \)
leads to a stationary vector of \( P = A + dr^T \), but also, in
a rather oblique manner, describe where \( x \) ends up if we start
UpdateIteration \((A, x, y)\) with an arbitrary \( y \).

**Theorem 1.** For vectors \( x, y, d \) and substochastic matrix
\( A \), if \( y = Ax - x + d \) and \( d \) is a non-negative vector, then
defining the stochastic matrix \( P = A + dr^T /\|d\|_1 \),

\[
\|Px - x\|_1 \leq \|y\|_1 \quad \text{and} \quad \|x\|_1 \geq \|d\|_1 - \|y\|_1.
\]

To reiterate, Theorem 1 not only describes the correct initial
value of \( y \) to arrive at a stationary vector of \( P = A + dr^T \),
but also says that for any \( A, x, y \), if the vector \( d \) satisfying
\( y = Ax - x + d \) is non-negative, then \( x \) arrives as the sta-
tionary distribution of a random walk on \( A \) that resets to a
distribution proportional to \( d \).

While the limit of \( x \) is well defined, choosing \( z \) arbitrarily
clearly need not result in rapid, or any, convergence to this
limit. Much as \( y \) leads \( x \) to its limit, vectors \( z \) whose coor-
dinates agree with those of \( y \) also exhibit brisk convergence
of \( \|y\|_1 \) to zero.

**Theorem 2.** If each \( z_u \) lies between zero and \( y_u \), then

\[
\|y + Az - z\|_1 \leq \|y\|_1 - \sum_u r_u|z_u|.
\]

When all \( r_u \) are equal, the exponential convergence of
PowerIteration is a special case of Theorem 2: processing
\( z = y \) each round reduces \( \|y\|_1 \) by a factor of \( 1 - r_u \). Mor-
over, when \( r \) is not uniform Theorem 2 gives a tighter char-
acterization of progress than eigenvalue bounds, which are
generally in terms of the smallest \( r_u \) value. Finally, and
critically, Theorem 2 describes progress made when we pro-
cess an update \( z \neq y \), and informs us as to where in \( y \) the
progress is being made.
3. ACCELERATION TECHNIQUES

In this section we consider several manners of choosing the vector \( z \) in UpdateIteration that give rise to various acceleration techniques. Most have occurred in some form previously in the literature, and we will discuss the often significant differences between their current and previous incarnations. In each case we will find shortcomings of previous techniques that are resolved by casting them in our common framework. Additionally, a significant advantage is the simple manner in which the techniques now compose, both from an algorithmic and performance perspective.

We also present experimental data detailing the performance of the acceleration techniques and compositions we discuss on a 34M page crawl from 2002 containing roughly 800M edges. The pages are organized first by host, where hosts are sorted by crawl discovery order, and within each host by crawl discovery order. Several benefits of such an ordering are discussed in Kamvar et al. [6], who use a more thorough sorting within each host. We choose to order pages by hosts, independent of the significant performance gains noted in [6], because one of our optimizations relies on it, and we require a consistent experimental framework.

For each approach, we plot the total error \( \|P\vec{x} - \vec{x}\|/\|\vec{x}\| \) against computational effort, measured in units of 800M edges processed, corresponding to the effort required by a single pass of PowerIteration. The normalization by \( \|\vec{x}\| \) is required because our vector \( \vec{x} \) need not remain at unit norm, and it would be unfair for a vector to achieve small \( \|P\vec{x} - \vec{x}\| \) simply by virtue of a small \( \vec{x} \). In reading the graphs, the acceleration can be seen by in the ratios of effort along a fixed (horizontal) level of error.

3.1 Sequential Updates

In choosing an update vector \( z \), each coordinate makes a commitment to the update \( z_u \) it intends, at which point each update is applied to \( x \) and propagated through \( A \) in parallel. However, in most implementations these updates will be processed serially, typically reading and propagating each \( z_u \) in turn. As the \( z_u \) may be chosen arbitrarily, there is no need for a node to commit to a particular value until it is needed. Rather, we can delay the choice of \( z_u \) until it is needed, conceptually processing a long series of single coordinate updates of the form \( z = (0, \ldots, 0, z_u, 0, \ldots) \).

Sequential updating allows us to base \( z_u \) on a value of \( y_u \) that reflects all updates applied thus far, even those applied in the current iteration, allowing us to propagate the effect of a single update multiple times in a single pass over the edge file. Even if the nodes are ordered randomly, roughly half of the edges will point forward in the node order. Updates passed along these edges will be processed before we complete a pass over the edge file. Well organized graphs can benefit even more, with updates pushed along entire acyclic subgraphs in one pass.

Sequential updating is based on a specific ordering of nodes, and clearly some orderings are better than others. The ordering we use is based on crawl order, which has the peculiar property that 80% of the edge point backwards; crawling very quickly discovers pages with many incoming links, and placing them early reverses the direction of the bulk of their links. As any ordering can easily be run in both orders, forwards or backwards or both, we will also consider the sequential updating in reverse order. Experimentally, alternating direction, forwards then backwards, performs more poorly than either unidirectional approach. This is peculiar, and merits further investigation.

Figures 1 and 2 compare traditional PowerIteration (PI) against sequential updating (SU) and sequential updating applied in reverse order (R-SU). It should be stressed that these techniques exhibit exactly the same data access patterns as traditional PowerIteration, passing linearly over the edge file and probing \( u \) in main memory for each edge \( A_{uv} \). The approaches differ only in what they do for each \( A_{uv} \) (and the direction of scan, for R-SU). Their running times are effectively identical to PowerIteration.

![Figure 1: Sequential Updates: Total Error](image1.png)

We see acceleration of nearly 2x and 3x for sequential updates on the crawl ordered and reverse crawl ordered graphs, respectively, with the gap between SU and R-SU diminishing with time.

![Figure 2: Sequential Updates: Maximum Error](image2.png)

Here we see again a lead of reverse crawl ordering, but the lead is less initially, growing after several iterations.

**Related Work:** Sequential updating is similar to the Gauss-Seidel approach described by Arasu et al. [1], in which one sequentially sets \( x_u = \sum P_{uv}x_v \), using the most current values of \( x_v \) rather than those of the previous iteration. In contrast with UpdateIteration, the Gauss-Seidel approach requires the graph’s edges to be grouped by destination, rather than source, which can substantially complicate data maintenance.
3.2 Reiterated Updates

A large fraction of links in the web graph are *intra-host*, and as such it is common to group pages by host for locality benefits, discussed in [6]. Given such a grouping, after processing the nodes associated with a host, a large fraction of the propagated update \( z \) will return to nodes on that host. While this may seem frustrating at first, recall that Theorem 2 says that \( |y||z| \) decreases by at least \( r_{u}|z_u| \), independent of where \( Az \) ends up. Moreover, various caches will retain the data used to process this group, making re-processing it very efficient. Rather than process the next group, reading sequential edge data from disk and probing \( y_v \) in main memory, we can reprocess the current group, reading sequential edge data from main memory and probing \( y_v \) in the L3 cache. The latter is substantially faster than the former, and represents a good payoff so long as substantial updates remain in the group. As the intra-host edge density is high, we might perform several iterations on a group before its \( y \) updates dissipate to other groups.

Another popular grouping is by strongly connected component. Ordered topologically, there is no reason to advance from a component until it has satisfactorily converged, as there are no edges along which updates from subsequent components may return. This approach has the decided advantage that the working set of edges and nodes at any point in time is only as large as the associated strongly connected component, each of which is visited only once. The main disadvantage is that computing strongly connected components is difficult in external memory, and an approximation should probably be used instead. Notice that we do not actually require that the grouping have no back edges, but the fewer that exist, the fewer updates return upstream and the more effective each pass is. Strongly connected components ensure that one pass suffices, but groupings that simply have low reverse edge density are highly effective as well.

There are other interesting groupings that one can imagine (we will discuss some more in Sections 3.3 and 3.4) and the question quickly emerges of which one should be used. In fact, we can use several. Our main constraint is that we access the edge file sequentially, and therefore we must collocate edges from nodes in the same group. If we have the disk space to maintain multiple edge files, we can produce an edge file for each grouping, and choose to use a particular edge file based on our needs at the time. In reading the edge file, we process the collocated nodes in a group, and can easily make multiple passes over this data without tripping over the intervening edges in the original edge file. This approach does not give us locality of reference in the \( y \) vector, as we have not actually changed node indices.

Figures 3 and 4 examine the performance benefits of grouping by host and processing each one, two, and three times. We also examine 10x reiteration, though only to demonstrate its limit. As we count operations instead of measure execution time, we will need to make some assumptions about the execution time of subsequent passes. For presentation reasons, we will assume that subsequent iterations are free, which is clearly false. However, the 2x and 3x reiterations result in acceleration of nearly 2x and 3x, respectively, so acceleration clearly exists for more pessimistic assumptions. Actual runtimes suggest that subsequent iterations are cheap, with 2x and 3x reiteration taking roughly 1.25 and 1.50 times as long, respectively, in a not especially well controlled environment. Also, we only need to process the intra-group edges while reiterating, propagating updates along inter-group edges only once finished.

The acceleration for total error is almost 2x and 3x over SU, suggesting that reiterated updates can be nearly as effective as multiple passes over the matrix. Of course, there are diminishing returns, visible as 3x and 10x converge.

Figures 3 and 4 examine the performance benefits of grouping by host and processing each one, two, and three times. We also examine 10x reiteration, though only to demonstrate its limit. As we count operations instead of measure execution time, we will need to make some assumptions about the execution time of subsequent passes. For presentation reasons, we will assume that subsequent iterations are free, which is clearly false. However, the 2x and 3x reiterations result in acceleration of nearly 2x and 3x, respectively, so acceleration clearly exists for more pessimistic assumptions. Actual runtimes suggest that subsequent iterations are cheap, with 2x and 3x reiteration taking roughly 1.25 and 1.50 times as long, respectively, in a not especially well controlled environment. Also, we only need to process the intra-group edges while reiterating, propagating updates along inter-group edges only once finished.

**Figure 3: Reiteration: Total Error**

Additional iterations help maximum error as well, but the diminishing returns are even more pronounced.

**Related Work:** Arrangement by strongly connected components has appeared several times in various forms. Eiron et al. [4] note that pages with no outlinks form a large fraction of the web and describe how to infer their ranks from the stationary probabilities of a modified graph with these nodes removed. More generally, Arasu et al. [1] and Langville and Meyer [10] view the problem as a block upper triangular linear system, processing strongly connected components in turn. Their techniques focus on decomposing the Markov chain, and require a strict topological order.

These approaches are captured by reiteration over the equivalent grouping. Moreover, UpdateIteration can take advantage of groupings that are only mostly topological. This flexibility addresses concerns of the substantial effort needed for data organization and maintenance, and enables grouping by host/domain, which was not possible in the more rigid block triangular techniques.
3.3 Selective Updates

While \( z_u = y_u \) is clearly one effective choice, an alternate choice is \( z_u = 0 \). In effect, we can choose not to update node \( u \). Clearly if \( y_u = 0 \) we need not expend effort to propagate \( y_u \) through \( A \), as we will simply be adding zero to several locations in \( y \). Even when \( y_u \) is non-zero but small, we may want to defer the update until the gains are more in line with the typical entry. With this in mind, there are various predicates we could use to decide if we should process \( y_u \).

We will specifically consider:

\[
\text{Effort} : \quad \text{Set} \ z_u = y_u \quad \text{iff} \quad \frac{|r_u y_u|}{\deg_u} \geq \text{avg} \left\{ \frac{|r_v y_v|}{\deg_v} \right\},
\]

selecting entries with the highest anticipated progress \(|r_u y_u|/\deg_u\) per expended effort \(\deg_u\). There are other predicates that could be used, each resulting from a different view of which entries are important to process. Examples include choosing those entries with largest relative error \(|(P x - x)_u|/|x_u|\) or those entries whose range of possible ordinal ranks is largest.

Selective updating has some interesting interaction with sequential and reiterated updates. As we run a pass of sequential updates the average value of \(|r_u y_u|/\deg_u\) will change, and while we could maintain the average exactly by carefully watching the changes in \( y \), we can also do a more efficient approximation by assuming that \(|y|_1\) decreases by exactly \( r_u|z_u| \). Reiteration is similar, in that each reiteration lowers the weight in a group markedly, by a factor of at least 1 − \( r_u \), but not the average value over all of \( y \). We could base our decision on the group’s average, shrinking with the values we consider, or on the entire average over \( y \).

While the gains of selective updating in terms of computation and memory accesses are clear, savings in terms of disk accesses are less so. It is not possible to skip entries on disk at no cost; data is read from disk in blocks, and the cost is amortized over all entries in the block. Likewise, disk prefetching will prepare subsequent blocks cheaply, and it is unclear that we gain anything by ignoring edge data passed to us. To address this somewhat, it is certainly possible to apply selective updating at a coarser scale than the node level. One could skip entire groups of entries at a time, permitting a volume of edges to be passed over at once and resulting entire disk blocks skipped.

Alternatively, Kamvar, Haveliwala, and Golub note in [5] that some pages converge more slowly than others, determining which these are at runtime by observing their relative change in ranks each round. While they use converged values to cull edges associated with converged nodes, we might base a grouping scheme (a la Section 3.2) on convergence rate, determined in a similar manner at run time. Emitting an appropriately grouped edge file can be done efficiently in a single pass so long as the number of groups is not terribly large. This organization of the edge file allows efficient passes over prefixes of the edge file, letting us efficiently process each group at an arbitrary rate. Understanding and experimenting with coarse-grained selective updating and grouping is interesting future research.

Figures 5 and 6 examine the Effort predicate applied to previous schemes (denoted in the figures labels by “E+”). The acceleration we see here is substantial, as selective updating takes advantage of the initially high variability of magnitudes in \( y \). We stress that actual acceleration will be less, although some may be recouped via clever grouping.

![Figure 5: Selective Updating: Total Error](image5)

We see substantial acceleration in terms of edges processed, which is, admittedly, a somewhat suspect measure. The message is that the work that needs to be done is less.

![Figure 6: Selective Updating: Maximum Error](image6)

Initial acceleration is especially pronounced here, as the selective updates immediately leap two orders of magnitude.

Related Work: Selective updating can be seen in the work of Kamvar, Haveliwala, and Golub [5], who describe a power iteration process wherein entries \( x_u \) that appear to have converged are frozen, saving them the recomputation of these entries every iteration. Moreover, edges associated with the frozen nodes are trimmed from the edge file, reducing disk IO required. Freezing is analogous to the non-transmission of an update, and the edge trimming is clearly the basis of the grouping discussed previously, though more final.

The main difference between [5] and UpdateIteration is that in the former the recipient decides whether an update will be propagated, and it is conceivable that significant updates may be ignored. This is particularly evident during incremental changes to the web graph. As the matrix \( A \) changes over time, previous stationary distributions prove good starting points for converging to the new stationary distribution. But if only a few links change, entries of \( x_u \) not incident to a changed edge will remain stationary after an iteration and will be frozen and never updated.
3.4 Incremental Updates

Over time the adjacency structure of the web changes, and we will want to compute the stationary distribution of a new chain that differs from the old in a relatively small number of locations. The stationary distribution of the old chain is generally viewed as a good first approximation, and indeed it is easy and intelligent to restart PowerIteration on a new chain using the old z.

Restarting UpdateIteration from a specific x appears non-trivial, as we must compute \( y = Ax - x + d \), involving a matrix multiplication. In fact, the process is much simpler: Let \( A \) and \( B \) describe the old and new edge transition matrices, and consider the two associated update vectors \( y_A \) and \( y_B \):

\[
y_A = Ax - x + d \quad \text{and} \quad y_B = Bx - x + d.
\]

We can relate the update vector \( y_B \) to its antecedent \( y_A \) as

\[
y_B = y_A + (B - A)x.
\]

This equivalence shows how to efficiently update \( y_A \) to \( y_B \), allowing us to efficiently reinvoke UpdateIteration\((B, x, y_A)\).

The effort required in this matrix-vector multiplication is proportional to the number of non-zero entries in \( B - A \), corresponding to the number of changed edge weights.

Several approaches to personalization of PageRank are based on personalization of the reset distribution [8, 9], shifting influence to those sites that the distribution favors, and the site linked by them. Recall from Theorem 1 that \( x \) converges to the stationary distribution of the chain with reset distribution \( d = y - (Ax - x) \). Personalization of the reset distribution is easily performed by incorporating any changes to \( d \) into \( y \) instead, changing \( x \)'s limit appropriately. It is worth stressing that \( x \)'s limit is defined by the distribution proportional to \( d \), and we need not worry about renormalizing \( d \) if we only make a few changes.

Finally, much of research into Markov chain based ranking research is exploratory: the best setting of weights in \( A \) and vectors \( d \) and \( r \) are not known. Uniform weights seem natural as defaults, but are clearly primitive choices. Exploring link weighting schemes based on content analysis or resetting policies based on content quality require efficient recomputation of ranks. Each of these explorative choices: updating \( A_{vu} \), \( r_u \), and \( d_u \) values, is easy in UpdateIteration, corresponding to simple updates to \( y \).

In these three cases above, the changes to the Markov chain often result in sparse updates to \( y \): most of the edges in the graph are stable between recrawls, and much of personalization of reset distributions is localized (upweighting a few trusted/bookmarked pages, for example). In this context, selective updating of Section 3.3 is well suited to efficiently process just those substantial entries, and leave the converged regions of the graph untouched. Of course to accommodate this properly, it makes sense to maintain an edge file of these parts of the graph that experience frequent edge churn, so that we needn't pass over the entire graph.

The fine granularity of sequential updates also allows a very smooth incremental update: we can decompose any update to the adjacency matrix into a set of small updates to the links of each node, which we apply as we visit each node. We need not pause the system to compute \((B - A)x\), but can apply the implications of changes at each node in turn. This becomes all the more relevant in a distributed setting where such pauses could destroy parallelism.

Figure 7 compares various techniques applied to a converged vector \( y \) that has had 1000 random positions updated randomly by \( \pm 1/n \), emulating either a change in the link structure or reset distribution. For small initial \( \|y\|_1 \), the scale of the updates does not affect the shape of the curves, so the choice \( 1/n \) is arbitrary.

![Figure 7: Incremental Updating](image-url)

It is difficult to characterize the acceleration of the incremental updates by a multiplicative factor, as it is clearly a different shape than the standard curves. Several orders of magnitude are gained immediately, with the slope arriving at the shape of Figure 5 as the initially concentrated \( y \) vector is distributed more uniformly.

![Figure 8: Incremental Updating](image-url)

Maximum error exhibits the same behavior as total error, dropping rapidly as the initially sparse vector is dispersed. The initial hiccup again reflects the sensitive nature of the maximum error measure.

Related Work: Chien et al. [3] describe an approach to incremental updating that is based on the construction and analysis of a new a Markov chain on nodes within a modest neighborhood of the graph changes, and a supernode representing nodes outside this horizon. Their approach is similar in spirit to ours, in that attention is restricted to the relatively small region where change may occur. However, rather than fix a region and degree of accuracy, UpdateIteration discovers where updates are needed as it goes, accommodating any degree of accuracy fluidly.
Haveliwala [8] and Jeh and Widom [9] have done work on efficient personalization, observing that the function mapping reset distributions to stationary distributions is linear. This enables very efficient manners of synthesizing personalized PageRanks from a set of precomputed PageRanks based on various reset distributions. For example, a page’s $d_u$ value can be increased by folding in the stationary distribution of a random walk that resets to only that page, exactly analogous to increasing and propagating $y_u$.

### 3.5 Floating Point Implications

Our ability to choose $z$ arbitrarily has implications for floating point error. We have the flexibility to always choose $z_u$ to be a power of two, so that its addition to $x$ will result in nominal floating point loss. This is harder to guarantee with $y$, as transmission along weighted edges will change $z_u$ from a power of two. Understanding and improving floating point behavior has positive implications for the introduction of strength reduction and low precision arithmetic, of particular interest in this setting where maintaining all of $x$ or $y$ in memory is challenging. Additionally, UpdateIteration propagates and combines updates $z_u$, which are typically of smaller magnitude than the $x_u$ values that PowerIteration operates with, and the precision maintained is thus higher.

### 3.6 Distribution and Robustness

If we remove the sequential behavior from sequential updates, we see that updates in UpdateIteration can actually be totally asynchronous. Moreover, our choices for $z_u$ are made locally with only a modicum of global information. This allows for a very smooth distributed implementation, in which the only coordination between compute nodes that is required is eventual communication of the updates applied. We can delay and reorder inter-node $z$ transmissions until the updates are significant, batching and trimming network overhead. Clearly, the best update schedule is highly dependent on the system topology, and we refrain from giving explicit suggestions here.

In an extreme case of delay, a compute node may be unavailable for a long period of time or even crash. Other compute nodes can continue in its absence, functioning under the belief that the PageRanks associated with that compute node simply have not changed. If the node comes online again it simply reenters the computation, transmitting and receiving updates. As noted for incremental updates, the granularity of sequential updating is very fine, and the amount of work needed to roll forward from any checkpoint can be arbitrarily small.

### 3.7 Decentralization

The Markov chain we have studied simulates the propagation of probability mass through a directed communication network whose nodes happen to be computational agents. The propagation of updates is easily performed within the communication network, as updates are only transmitted along links. The initial values of $x = 0$ and $y = d$ are easily chosen, as $d$ needn’t be normalized. As the network changes, in the incremental fashion suggested by Section 3.4, the necessary updates to $y$ are computable by the source of the edge that has arrived or departed. Gracefully departing nodes removing their incoming edges using this update mechanism and apply the update $z_u = y_u - d_u$ before departing.

### 4. CONCLUSIONS AND FUTURE WORK

We have examined an algorithmic reformulation of the traditional power iteration algorithm based on the propagation of updates rather than values. UpdateIteration enables several algorithmic optimizations that result in more efficient convergence. Moreover, the optimizations are well suited to the problems of incremental and personalized updates to the underlying Markov chain, and permit flexible operation in a distributed setting.

The optimizations presented here are likely just a sampling of what can be done to accelerate computation of PageRank. These optimizations are intended to take advantage of particular features of computer systems, and it seems likely that other features may yet be exploited, both for performance and potentially quality of ranking. Techniques such as Arnoldi Iteration and unsymmetric Lanczos are tempting targets, as is the power extrapolation approach of Kamvar et al. [7]. Additionally, there is work to be node exploring the new possibilities enabled through efficient PageRank computation.

### 5. ACKNOWLEDGMENTS

The author would like to thank several people who contributed constructive ideas and observations. Michael Isard, Steve Chien, Kevin McCurley, the participants of the Workshop on Search and Meta-Search, and the anonymous reviewers all gave valuable comments which have greatly improved the presentation.

### 6. REFERENCES


7. APPENDIX A: PROOFS

We now look at the two deferred proofs from Section 2. Recall that Theorem 1 requires the entries of $d$ be non-negative.

Proof of Theorem 1. $Px - x$ and $y = Ax - x + d$ differ only in the amount of $d$ added to $Ax - x$. We can thus write their difference as

$$y - (Px - x) = d - dr^T x/∥d∥_1.$$ (1)

Summing the coordinates of vectors on both sides of (1), and noting that $\sum_u (Px)_u = \sum_u x_u$ and $\sum_u d_u = ∥d∥_1$, gives

$$\sum_u y_u = ∥d∥_1 - r^T x .$$ (2)

To prove the first stated inequality, we move $y$ to the right hand side of (1), take norms, and use the triangle inequality.

$$∥Px - x∥_1 ≤ ∥y∥_1 + |(∥d∥_1 - r^T x)| .$$ (3)

Substituting $\sum_u y_u$ for $∥d∥_1 - r^T x$ and then $| \sum_u y_u | ≤ ∥y∥_1$, we get

$$∥Px - x∥_1 ≤ ∥y∥_1 + | \sum_u y_u | ≤ 2∥y∥_1 .$$ (4)

Similarly, the second stated inequality results from the inequalities

$$∥x∥_1 ≥ r^T x = ∥d∥_1 - \sum_u y_u ≥ ∥d∥_1 - ∥y∥_1 .$$ (5)

with the inequality $∥x∥_1 ≥ r^T x$ following as all $|r_u| ≤ 1 . □$

The proof of Theorem 2 relies on the assumption that the coordinates of $z$ lie between zero and the corresponding $y_u$.

Proof of Theorem 2. As each $z_u$ lies between zero and $y_u$, we have that $∥y - z∥_1 = ∥y∥_1 - ∥z∥_1$, and starting from the triangle inequality

$$∥y + Az - z∥_1 ≤ ∥y - z∥_1 + ∥Az∥_1$$ (6)

we get

$$∥y + Az - z∥_1 = ∥y∥_1 - ∥z∥_1 + ∥Az∥_1 .$$ (7)

Column $u$ of $A$ sums to $r_u$, and thus $∥Az∥_1 ≤ \sum_u (1-r_u)|z_u|$. We thus get

$$∥y + Az - z∥_1 ≤ ∥y∥_1 - \sum_u |z_u| + \sum_u (1-r_u)|z_u| .$$ (8)

Collecting the summands yields the claimed bound. □