Bayesian Compressive Sensing Via Belief Propagation

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Abstract—Compressive sensing (CS) is an emerging field based on the revelation that a small collection of linear projections of a sparse signal contains enough information for stable, sub-Nyquist signal acquisition. When a statistical characterization of the signal is available, Bayesian inference can complement conventional CS methods based on linear programming or greedy algorithms. We perform asymptotically optimal Bayesian inference using belief propagation (BP) decoding, which represents the CS encoding matrix as a graphical model. Fast computation is obtained by reducing the size of the graphical model with sparse encoding matrices. To decode a length-N signal containing K large coefficients, our CS-BP decoding algorithm uses $O(K \log(N))$ measurements and $O(N \log^2(N))$ computation. Finally, although we focus on a two-state mixture Gaussian model, CS-BP is easily adapted to other signal models.

Index Terms—Bayesian inference, belief propagation, compressive sensing, fast algorithms, sparse matrices.

I. INTRODUCTION

ANY signal processing applications require the identification and estimation of a few significant coefficients from a high-dimensional vector. The wisdom behind this is the ubiquitous compressibility of signals: in an appropriate basis, most of the information contained in a signal often resides in just a few large coefficients. Traditional sensing and processing first acquires the entire data, only to later throw away most coefficients and retain the few significant ones [2]. Interestingly, the information contained in the few large coefficients can be captured (encoded) by a small number of random linear projections [3]. The ground-breaking work in compressive sensing (CS) [4]–[6] has proved for a variety of settings that the signal can then be decoded in a computationally feasible manner from these random projections.

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A. Compressive Sensing

Sparsity and Random Encoding: In a typical compressive sensing (CS) setup, a signal vector $x \in \mathbb{R}^N$ has the form $x = \Psi \theta$, where $\Psi \in \mathbb{R}^{N \times N}$ is an orthonormal basis, and $\theta \in \mathbb{R}^N$ satisfies $\|\theta\|_0 = K \ll N.^1$ Owing to the sparsity of x relative to the basis Ψ , there is no need to sample all N values of x. Instead, the CS theory establishes that x can be decoded from a small number of projections onto an incoherent set of measurement vectors [4], [5]. To measure (encode) x, we compute $M \ll N$ linear projections of x via the matrix-vector multiplication $y = \Phi x$ where $\Phi \in \mathbb{R}^{M \times N}$ is the encoding matrix.

In addition to *strictly sparse* signals where $||\theta||_0 \le K$, other *signal models* are possible. *Approximately sparse* signals have $K \ll N$ large coefficients, while the remaining coefficients are small but not necessarily zero. *Compressible* signals have coefficients that, when sorted, decay quickly according to a power law. Similarly, both *noiseless* and *noisy* signals and measurements may be considered. We emphasize noiseless measurement of approximately sparse signals in the paper.

Decoding via Sparsity: Our goal is to decode x given y and Φ . Although decoding x from $y = \Phi x$ appears to be an ill-posed inverse problem, the prior knowledge of sparsity in x enables to decode x from $M \ll N$ measurements. Decoding often relies on an optimization, which searches for the sparsest coefficients θ that agree with the measurements y. If M is sufficiently large and θ is strictly sparse, then θ is the solution to the ℓ_0 minimization:

$$\hat{\theta} = \arg\min ||\theta||_0 \text{ s.t. } y = \Phi \Psi \theta.$$

Unfortunately, solving this ℓ_0 optimization is NP-complete [7]. The revelation that supports the CS theory is that a computationally tractable optimization problem yields an equivalent solution. We need only solve for the ℓ_1 -sparsest coefficients that agree with the measurements y [4], [5]

$$\hat{\theta} = \arg\min \|\theta\|_1 \text{ s.t. } y = \Phi \Psi \theta$$
 (1)

as long as $\Phi\Psi$ satisfies some technical conditions, which are satisfied with overwhelming probability when the entries of Φ are independent and identically distributed (i.i.d.) sub-Gaussian random variables [4]. This ℓ_1 optimization problem (1), also known as *Basis Pursuit* [8], can be solved with linear programming methods. The ℓ_1 decoder requires only $M = O(K \log(N/K))$ projections [9], [10]. However,

 $^1\mbox{We use}\parallel\cdot\parallel_0$ to denote the ℓ_0 "norm" that counts the number of nonzero elements.

encoding by a dense Gaussian Φ is slow, and ℓ_1 decoding requires cubic computation in general [11].

B. Fast CS Decoding

While ℓ_1 decoders figure prominently in the CS literature, their cubic complexity still renders them impractical for many applications. For example, current digital cameras acquire images with $N=10^6$ pixels or more, and fast decoding is critical. The slowness of ℓ_1 decoding has motivated a flurry of research into faster algorithms.

One line of research involves iterative greedy algorithms. The Matching Pursuit (MP) [12] algorithm, for example, iteratively selects the vectors from the matrix $\Phi\Psi$ that contain most of the energy of the measurement vector y. MP has been proven to successfully decode the acquired signal with high probability [12], [13]. Algorithms inspired by MP include OMP [12], tree matching pursuit [14], stagewise OMP [15], CoSaMP [16], IHT [17], and Subspace Pursuit [18] have been shown to attain similar guarantees to those of their optimization-based counterparts [19]–[21].

While the CS algorithms discussed above typically use a dense Φ matrix, a class of methods has emerged that employ *structured* Φ . For example, subsampling an orthogonal basis that admits a fast implicit algorithm also leads to fast decoding [4]. Encoding matrices that are themselves sparse can also be used. Cormode and Muthukrishnan proposed fast streaming algorithms based on group testing [22], [23], which considers subsets of signal coefficients in which we expect at most one "heavy hitter" coefficient to lie. Gilbert *et al.* [24] propose the Chaining Pursuit algorithm, which works best for extremely sparse signals.

C. Bayesian CS

CS decoding algorithms rely on the sparsity of the signal x. In some applications, a statistical characterization of the signal is available, and Bayesian inference offers the potential for more precise estimation of x or a reduction in the number of CS measurements. Ji et al. [25] have proposed a Bayesian CS framework where relevance vector machines are used for signal estimation. For certain types of hierarchical priors, their method can approximate the posterior density of x and is somewhat faster than ℓ_1 decoding. Seeger and Nickisch [26] extend these ideas to experimental design, where the encoding matrix is designed sequentially based on previous measurements. Another Bayesian approach by Schniter et al. [27] approximates conditional expectation by extending the maximal likelihood approach to a weighted mixture of the most likely models. There are also many related results on application of Bayesian methods to sparse inverse problems (cf. [28] and references therein).

Bayesian approaches have also been used for multiuser decoding (MUD) in communications. In MUD, users modulate their symbols with different spreading sequences, and the received signals are superpositions of sequences. Because most users are inactive, MUD algorithms extract information from a sparse superposition in a manner analogous to CS decoding.

Guo and Wang [29] perform MUD using sparse spreading sequences and decode via belief propagation (BP) [30]–[35]; our paper also uses sparse encoding matrices and BP decoding. A related algorithm for decoding low density lattice codes (LDLC) by Sommer *et al.* [36] uses BP on a factor graph whose self and edge potentials are Gaussian mixtures. Convergence results for the LDLC decoding algorithm have been derived for Gaussian noise [36].

D. Contributions

In this paper, we develop a sparse encoder matrix Φ and a belief propagation (BP) decoder to accelerate CS encoding and decoding under the Bayesian framework. We call our algorithm CS-BP. Although we emphasize a two-state mixture Gaussian model as a prior for sparse signals, CS-BP is flexible to variations in the signal and measurement models.

Encoding by Sparse CS Matrix: The dense sub-Gaussian CS encoding matrices [4], [5] are reminiscent of Shannon's random code constructions. However, although dense matrices capture the information content of sparse signals, they may not be amenable to fast encoding and decoding. Low density parity check (LDPC) codes [37], [38] offer an important insight: encoding and decoding are fast, because multiplication by a sparse matrix is fast; nonetheless, LDPC codes achieve rates close to the Shannon limit. Indeed, in a previous paper [39], we used an LDPC-like sparse Φ for the special case of noiseless measurement of strictly sparse signals; similar matrices were also proposed for CS by Berinde and Indyk [40]. Although LDPC decoding algorithms may not have provable convergence, the recent extension of LDPC to LDLC codes [36] offers provable convergence. Additionally, CS-BP has recently been proved to be asymptotically optimal in the large system limit [29], [60].

We encode (measure) the signal using sparse Rademacher $(\{0,1,-1\})$ LDPC-like Φ matrices. Because entries of Φ are restricted to $\{0,1,-1\}$, encoding only requires sums and differences of small subsets of coefficient values of x. The design of Φ , including characteristics such as column and row weights, is based on the relevant signal and measurement models, as well as the accompanying decoding algorithm.

Decoding by BP: We represent the sparse Φ as a sparse bipartite graph. In addition to accelerating the algorithm, the sparse structure reduces the number of loops in the graph and thus assists the convergence of a message passing method that solves a Bayesian inference problem. Our estimate for x explains the measurements while offering the best match to the prior. We employ BP in a manner similar to LDPC channel decoding [34], [37], [38]. To decode a length-N signal containing K large coefficients, our CS-BP decoding algorithm uses $M = O(K \log(N))$ measurements and $O(N \log^2(N))$ computation. Although CS-BP is not guaranteed to converge, numerical results are quite favorable.

The remainder of the paper is organized as follows. Section II defines our signal model, and Section III describes our sparse CS-LDPC encoding matrix. The CS-BP decoding algorithm is described in Section IV, and its performance is demonstrated

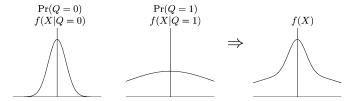


Fig. 1. Mixture Gaussian model for signal coefficients. The distribution of X conditioned on the two state variables, Q=0 and Q=1, is depicted. Also shown is the overall distribution for X.

numerically in Section V. Variations and applications are discussed in Section VI, and Section VII concludes.

II. MIXTURE GAUSSIAN SIGNAL MODEL

We focus on a two-state mixture Gaussian model [41]–[43] as a prior that succinctly captures our prior knowledge about approximate sparsity of the signal. Bayesian inference using a two-state mixture model has been studied well before the advent of CS, for example by George and McCulloch [44] and Geweke [45]; the model was proposed for CS in [1] and also used by He and Carin [46]. More formally, let $X = [X(1), \dots, X(N)]$ be a random vector in \mathbb{R}^N , and consider the signal x = [x(1), ..., x(N)] as an outcome of X. Because our approximately sparse signal consists of a small number of large coefficients and a large number of small coefficients, we associate each probability density function (pdf) f(X(i)) with a state variable Q(i) that can take on two values. Large and small magnitudes correspond to zero mean Gaussian distributions with high and low variances, which are implied by Q(i) = 1 and Q(i) = 0, respectively,

$$\begin{split} f(X(i) \,|\, Q(i) = 1) &\sim \mathcal{N}\left(0, \sigma_1^2\right) \quad \text{and} \\ f(X(i) \,|\, Q(i) = 0) &\sim \mathcal{N}\left(0, \sigma_0^2\right), \end{split}$$

with $\sigma_1^2 > \sigma_0^2$. Let $Q = [Q(1), \dots, Q(N)]$ be the state random vector associated with the signal; the actual configuration $q = [q(1), \dots, q(N)] \in \{0,1\}^N$ is one of 2^N possible outcomes. We assume that the Q(i)'s are i.i.d.² To ensure that we have approximately K large coefficients, we choose the probability mass function (pmf) of the state variable Q(i) to be Bernoulli with $\Pr(Q(i) = 1) = S$ and $\Pr(Q(i) = 0) = 1 - S$, where S = K/N is the *sparsity rate*.

The resulting model for signal coefficients is a two-state mixture Gaussian distribution, as illustrated in Fig. 1. This mixture model is completely characterized by three parameters: the sparsity rate S and the variances σ_0^2 and σ_1^2 of the Gaussian pdf's corresponding to each state.

Mixture Gaussian models have been successfully employed in image processing and inference problems, because they are simple yet effective in modeling real-world signals [41]–[43]. Theoretical connections have also been made between wavelet coefficient mixture models and the fundamental parameters of Besov spaces, which have proved invaluable for characterizing real-world images. Moreover, arbitrary densities with a finite number of discontinuities can be approximated arbitrarily

²The model can be extended to capture dependencies between coefficients, as suggested by Ji *et al.* [25].

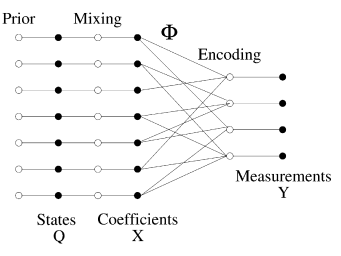


Fig. 2. Factor graph depicting the relationship between variable nodes (black) and constraint nodes (white) in CS-BP.

closely by increasing the number of states and allowing nonzero means [47]. We leave these extensions for future work, and focus on two-state mixture Gaussian distributions for modeling the signal coefficients.

III. SPARSE ENCODING

Sparse CS Encoding Matrix: We use a sparse Φ matrix to accelerate both CS encoding and decoding. Our CS encoding matrices are dominated by zero entries, with a small number of nonzeros in each row and each column. We focus on CS-LDPC matrices whose nonzero entries are $\{-1,1\}^3$; each measurement involves only sums and differences of a small subset of coefficients of x. Although the coherence between a sparse Φ and Ψ , which is the maximal inner product between rows of Φ and Ψ , may be higher than the coherence using a dense Φ matrix [48], as long as Φ is not too sparse (see Theorem 1 below) the measurements capture enough information about x to decode the signal. A CS-LDPC Φ can be represented as a bipartite graph G, which is also sparse. Each edge of G connects a coefficient node x(i) to an encoding node y(j) and corresponds to a nonzero entry of Φ (Fig. 2).

In addition to the core structure of Φ , we can introduce other constraints to tailor the measurement process to the signal model. The *constant row weight* constraint makes sure that each row of Φ contains exactly L nonzero entries. The row weight L can be chosen based on signal properties such as sparsity, possible measurement noise, and details of the decoding process. Another option is to use a *constant column weight* constraint, which fixes the number of nonzero entries in each column of Φ to be a constant R.

Although our emphasis is on noiseless measurement of approximately sparse signals, we briefly discuss noisy measurement of a strictly sparse signal, and show that a constant row weight L ensures that the measurements are approximated by two-state mixture Gaussians. To see this, consider a strictly

³CS-LDPC matrices are slightly different from LDPC parity check matrices, which only contain the binary entries 0 and 1. We have observed numerically that allowing negative entries offers improved performance. At the expense of additional computation, further minor improvement can be attained using sparse matrices with Gaussian nonzero entries.

sparse x with sparsity rate S and Gaussian variance σ_1^2 . We now have $y=\Phi x+z$, where $z\sim \mathcal{N}(0,\sigma_Z^2)$ is additive white Gaussian noise (AWGN) with variance σ_Z^2 . In our approximately sparse setting, each row of Φ picks up $\approx L(1-S)$ small magnitude coefficients. If $L(1-S)\sigma_0^2\approx\sigma_Z^2$, then the few large coefficients will be obscured by similar noise artifacts.

Our definition of Φ relies on the implicit assumption that x is sparse in the canonical sparsifying basis, i.e., $\Psi=I$. In contrast, if x is sparse in some other basis Ψ , then more complicated encoding matrices may be necessary. We defer the discussion of these issues to Section VI, but emphasize that in many practical situations our methods can be extended to support the sparsifying basis Ψ in a computationally tractable manner.

Information Content of Sparsely Encoded Measurements: The sparsity of our CS-LDPC matrix may yield measurements y that contain less information about the signal x than a dense Gaussian Φ . The following theorem, whose proof appears in the Appendix, verifies that y retains enough information to decode x well. As long as $S = K/N = \Omega((\sigma_0/\sigma_1)^2)$, then $M = O(K \log(N))$ measurements are sufficient.

Theorem 1: Let x be a two-state mixture Gaussian signal with sparsity rate S=K/N and variances σ_0^2 and σ_1^2 , and let Φ be a CS-LDPC matrix with constant row weight $L=\eta(\ln(SN^{1+\gamma}))/S$, where $\eta,\gamma>0$. If

$$M = O\left(\frac{(1+2\eta^{-1})(1+\gamma)}{\mu^2} \left[2K + (N-K) \left(\frac{\sigma_0}{\sigma_1}\right)^2 \right] \log(N) \right)$$
(2)

then x can be decoded to \hat{x} such that $||x - \hat{x}||_{\infty} < \mu \sigma_1$ with probability $1 - 2N^{-\gamma}$.

The proof of Theorem 1 relies on a result by Wang et~al. ([49], Theorem 1). Their proof partitions Φ into M_2 submatrices of M_1 rows each, and estimates each $\hat{x_i}$ as a median of inner products with submatrices. The ℓ_{∞} performance guarantee relies on the union bound; a less stringent guarantee yields a reduction in M_2 . Moreover, L can be reduced if we increase the number of measurements accordingly. Based on numerical results, we propose the following modified values as rules of thumb:

$$L \approx S^{-1} = N/K$$
, $M = O(K \log(N))$, and $R = LM/N = O(\log(N))$. (3)

Noting that each measurement requires O(L) additions and subtractions, and using our rules of thumb for L and M (3), the computation required for encoding is $O(LM) = O(N \log(N))$, which is significantly lower than the $O(MN) = O(KN \log(N))$ required for dense Gaussian Φ . In addition to Theorem 1, recent results indicate that there is no loss of optimality with respect to any random matrix distribution when using a sparse CS-LDPC matrix [29], [60]; CS-BP is asymptotically optimal in the large system limit.

IV. CS-BP DECODING OF APPROXIMATELY SPARSE SIGNALS

Decoding approximately sparse random signals can be treated as a Bayesian inference problem. We observe the measurements $y=\Phi x$, where x is a mixture Gaussian signal. Our goal is to estimate x given Φ and y. Because the set of equations $y=\Phi x$ is under-determined, there are infinitely many

solutions. All solutions lie along a hyperplane of dimension N-M. We locate the solution within this hyperplane that best matches our prior signal model. Consider the minimum mean-square error (MMSE) and maximum *a posteriori* (MAP) estimates:

$$\begin{split} \hat{x}_{\mathrm{MMSE}} &= \arg\min_{x'} E ||X - x'||_2^2 \text{ s.t. } y = \Phi x', \\ \hat{x}_{\mathrm{MAP}} &= \arg\max_{x'} f(X = x') \text{ s.t. } y = \Phi x', \end{split}$$

where the expectation is taken over the prior distribution for X. The MMSE estimate can be expressed as the conditional mean, $\hat{x}_{\text{MMSE}} = E[X \mid Y = y]$, where $Y \in \mathbb{R}^M$ is the random vector that corresponds to the measurements. Although the precise computation of \hat{x}_{MMSE} may require the evaluation of 2^N terms, a close approximation to the MMSE estimate can be obtained using the (usually small) set of state configuration vectors q with dominant posterior probability [27]. Indeed, exact inference in graphical models is NP-hard [50], because of loops in the graph induced by Φ . However, the sparse structure of Φ reduces the number of loops and enables us to use low-complexity message-passing methods to estimate x approximately.

A. Decoding Algorithm

We now employ belief propagation (BP), an efficient method for solving inference problems by iteratively passing messages over graphical models [30]–[35]. Although BP has not been proved to converge, for graphs with few loops it often offers a good approximation to the solution to the MAP inference problem. BP relies on *factor graphs*, which enable fast computation of global multivariate functions by exploiting the way in which the global function factors into a product of simpler local functions, each of which depends on a subset of variables [51].

Factor Graph for CS-BP: The factor graph shown in Fig. 2 captures the relationship between the states q, the signal coefficients x, and the observed CS measurements y. The graph is bipartite and contains two types of vertices; all edges connect variable nodes (black) and constraint nodes (white). There are three types of variable nodes corresponding to state variables Q(i), coefficient variables X(i), and measurement variables Y(j). The factor graph also has three types of constraint nodes, which encapsulate the dependencies that their neighbors in the graph (variable nodes) are subjected to. First, prior constraint nodes impose the Bernoulli prior on state variables. Second, mixing constraint nodes impose the conditional distribution on coefficient variables given the state variables. Third, encoding constraint nodes impose the encoding matrix structure on measurement variables.

Message Passing: CS-BP approximates the marginal distributions of all coefficient and state variables in the factor graph, conditioned on the observed measurements Y, by passing messages between variable nodes and constraint nodes. Each message encodes the marginal distributions of a variable associated with one of the edges. Given the distributions $\Pr(Q(i) \mid Y = y)$ and $f(X(i) \mid Y = y)$, one can extract MAP and MMSE estimates for each coefficient.

Denote the message sent from a variable node v to one of its neighbors in the bipartite graph, a constraint node c, by $\mu_{v\longrightarrow c}(v)$; a message from c to v is denoted by $\mu_{c\longrightarrow v}(v)$.

The message $\mu_{v\longrightarrow c}(v)$ is updated by taking the product of all messages received by v on all other edges. The message $\mu_{c\longrightarrow v}(v)$ is computed in a similar manner, but the constraint associated with c is applied to the product and the result is marginalized. More formally,

$$\mu_{v \longrightarrow c}(v) = \prod_{u \in n(v) \setminus \{c\}} \mu_{u \longrightarrow v}(v) \tag{4}$$

$$\mu_{c \longrightarrow v}(v) = \sum_{\sim \{v\}} \left(\operatorname{con}(n(c)) \prod_{w \in n(c) \setminus \{v\}} \mu_{w \longrightarrow c}(w) \right)$$
 (5)

where n(v) and n(c) are sets of neighbors of v and c, respectively, $\operatorname{con}(n(c))$ is the constraint on the set of variable nodes n(c), and $\sim \{v\}$ is the set of neighbors of c excluding v. We interpret these 2 types of message processing as $\operatorname{multiplication}$ of beliefs at variable nodes (4) and $\operatorname{convolution}$ at constraint nodes (5). Finally, the marginal distribution f(v) for a given variable node is obtained from the product of all the most recent incoming messages along the edges connecting to that node,

$$f(v) = \prod_{u \in n(v)} \mu_u \longrightarrow v(v). \tag{6}$$

Based on the marginal distribution, various statistical characterizations can be computed, including MMSE, MAP, error bars, and so on.

We also need a method to encode beliefs. One method is to sample the relevant pdf's uniformly and then use the samples as messages. Another encoding method is to approximate the pdf by a mixture Gaussian with a given number of components, where mixture parameters are used as messages. These two methods offer different trade-offs between modeling flexibility and computational requirements; details appear in Sections IV-B and IV-C. We leave alternative methods such as particle filters and importance sampling for future research.

Protecting Against Loopy Graphs and Message Quantization Errors: BP converges to the exact conditional distribution in the ideal situation where the following conditions are met: i) the factor graph is cycle-free; and ii) messages are processed and propagated without errors. In CS-BP decoding, both conditions are violated. First, the factor graph is loopy—it contains cycles. Second, message encoding methods introduce errors. These nonidealities may lead CS-BP to converge to imprecise conditional distributions, or more critically, lead CS-BP to diverge [52]–[54]. To some extent these problems can be reduced by i) using CS-LDPC matrices, which have a relatively modest number of loops; and ii) carefully designing our message encoding methods (Sections IV-B and IV-C). We stabilize CS-BP against these nonidealities using message damped belief propagation (MDBP) [55], where messages are weighted averages between old and new estimates. Despite the damping, CS-BP is not guaranteed to converge, and yet the numerical results of Section V demonstrate that its performance is quite promising. We conclude with a prototype algorithm; Matlab code is available at http://dsp.rice.edu/CSBP.

CS-BP Decoding Algorithm:

1) **Initialization**: Initialize the iteration counter i=1. Set up data structures for factor graph messages $\mu_{v\longrightarrow c}(v)$ and

- $\mu_{c\longrightarrow v}(v)$. Initialize messages $\mu_{v\longrightarrow c}(v)$ from variable to constraint nodes with the signal prior.
- 2) **Convolution**: For each measurement $c=1,\ldots,M$, which corresponds to constraint node c, compute $\mu_{c\longrightarrow v}(v)$ via convolution (5) for all neighboring variable nodes n(c). If measurement noise is present, then convolve further with a noise prior. Apply damping methods such as MDBP [55] by weighting the new estimates from iteration i with estimates from previous iterations.
- 3) Multiplication: For each coefficient v = 1,..., N, which corresponds to a variable node v, compute μ_{v→c}(v) via multiplication (4) for all neighboring constraint nodes n(v). Apply damping methods as needed. If the iteration counter has yet to reach its maximal value, then go to Step 2.
- 4) **Output**: For each coefficient v = 1, ..., N, compute MMSE or MAP estimates (or alternative statistical characterizations) based on the marginal distribution f(v) (6). Output the requisite statistics.

B. Samples of the PDF as Messages

Having described main aspects of the CS-BP decoding algorithm, we now focus on the two message encoding methods, starting with samples. In this method, we sample the pdf and send the samples as messages. Multiplication of pdf's (4) corresponds to point-wise multiplication of messages; convolution (5) is computed efficiently in the frequency domain.⁴

The main advantage of using samples is flexibility to different prior distributions for the coefficients; for example, mixture Gaussian priors are easily supported. Additionally, both multiplication and convolution are computed efficiently. However, sampling has large memory requirements and introduces quantization errors that reduce precision and hamper the convergence of CS-BP [52]. Sampling also requires finer sampling for precise decoding; we propose to sample the pdf's with a spacing less than σ_0

We analyze the computational requirements of this method. Let each message be a vector of p samples. Each iteration performs multiplication at coefficient nodes (4) and convolution at constraint nodes (5). Outgoing messages are modified,

$$\mu_{v \longrightarrow c}(v) = \frac{\prod_{u \in n(v)} \mu_{u \longrightarrow v}(v)}{\mu_{c \longrightarrow v}(v)} \quad \text{and}$$

$$\mu_{c \longrightarrow v}(v) = \sum_{v \in \{v\}} \left(\cos(n(c)) \frac{\prod_{w \in n(c)} \mu_{w \longrightarrow c}(w)}{\mu_{v \longrightarrow c}(v)} \right) \quad (7)$$

where the denominators are nonzero, because mixture Gaussian pdf's are strictly positive. The modifications (7) reduce computation, because the numerators are computed once and then reused for all messages leaving the node being processed.

Assuming that the column weight R is fixed (Section III), the computation required for message processing at a variable node is O(Rp) per iteration, because we multiply R+1 vectors of length p. With O(N) variable nodes, each iteration requires

 $^4\mathrm{Fast}$ convolution via FFT has been used in LDPC decoding over $GF(2^q)$ using BP [34].

O(NRp) computation. For constraint nodes, we perform convolution in the frequency domain, and so the computational cost per node is $O(Lp\log(p))$. With O(M) constraint nodes, each iteration is $O(LMp\log(p))$. Accounting for both variable and constraint nodes, each iteration is $O(NRp + LMp\log(p)) = O(p\log(p)N\log(N))$, where we employ our rules of thumb for L, M, and R (3). To complete the computational analysis, we note first that we use $O(\log(N))$ CS-BP iterations, which is proportional to the diameter of the graph [56]. Second, sampling the pdf's with a spacing less than σ_0 , we choose $p = O(\sigma_1/\sigma_0)$ to support a maximal amplitude on the order of σ_1 . Therefore, our overall computation is $O(\sigma_1/\sigma_0\log(\sigma_1/\sigma_0)N\log^2(N))$, which scales as $O(N\log^2(N))$ when σ_0 and σ_1 are constant.

C. Mixture Gaussian Parameters as Messages

In this method, we approximate the pdf by a mixture Gaussian with a maximum number of components, and then send the mixture parameters as messages. For both multiplication (4) and convolution (5), the resulting number of components in the mixture is multiplicative in the number of constituent components. To keep the message representation tractable, we perform model reduction using the Iterative Pairwise Replacement Algorithm (IPRA) [57], where a sequence of mixture models is computed iteratively.

The advantage of using mixture Gaussians to encode pdf's is that the messages are short and hence consume little memory. This method works well for mixture Gaussian priors, but could be difficult to adapt to other priors. Model order reduction algorithms such as IPRA can be computationally expensive [57], and introduce errors in the messages, which impair the quality of the solution as well as the convergence of CS-BP [52].

Again, we analyze the computational requirements. Because it is impossible to undo the multiplication in (4) and (5), we cannot use the modified form (7). Let m be the maximum model order. Model order reduction using IPRA [57] requires $O(m^2R^2)$ computation per coefficient node per iteration. With O(N) coefficient nodes, each iteration is $O(m^2R^2N)$. Similarly, with O(M) constraint nodes, each iteration is $O(m^2L^2M)$. Accounting for $O(\log(N))$ CS-BP iterations, overall computation is $O(m^2[L^2M + R^2N]\log(N)) = O(m^2(N)/(S)\log^2(N))$.

D. Properties of CS-BP Decoding

We briefly describe several properties of CS-BP decoding. The *computational characteristics* of the two methods for encoding beliefs about conditional distributions were evaluated in Sections IV-B and IV-C. The *storage requirements* are mainly for message representation of the $LM = O(N\log(N))$ edges. For encoding with pdf samples, the message length is p, and so the storage requirement is $O(pN\log(N))$. For encoding with mixture Gaussian parameters, the message length is m, and so the storage requirement is $O(mN\log(N))$. Computational and storage requirements are summarized in Table I.

Several additional properties are now featured. First, we have $progressive\ decoding$; more measurements will improve the precision of the estimated posterior probabilities. Second, if we are only interested in an estimate of the state configuration vector q but not in the coefficient values, then less information must

TABLE I COMPUTATIONAL AND STORAGE REQUIREMENTS OF CS-BP DECODING

Messages	Parameter	Computation	Storage
Samples of pdf	$p = O(\sigma_1/\sigma_0)$ samples	$O\left(\frac{\sigma_1}{\sigma_0}\log\left(\frac{\sigma_1}{\sigma_0}\right)N\log^2(N)\right)$	$O(pN\log(N))$
Mixture Gaussians	m components	$O\left(m^2 \tfrac{N}{S} \log^2(N)\right)$	$O(mN\log(N))$

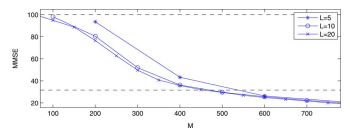


Fig. 3. MMSE as a function of the number of measurements M using different matrix row weights L. The dashed lines show the ℓ_2 norms of x (top) and the small coefficients (bottom). ($N=1000,S=0.1,\sigma_1=10,\sigma_0=1$, and noiseless measurements.)

be extracted from the measurements. Consequently, the number of measurements can be reduced. Third, we have *robustness to noise*, because noisy measurements can be incorporated into our model by convolving the noiseless version of the estimated pdf (5) at each encoding node with the pdf of the noise.

V. NUMERICAL RESULTS

To demonstrate the efficacy of CS-BP, we simulated several different settings. In our first setting, we considered decoding problems where $N = 1000, S = 0.1, \sigma_1 = 10, \sigma_0 = 1$, and the measurements are noiseless. We used samples of the pdf as messages, where each message consisted of $p = 525 = 3.5^2.7$ samples; this choice of p provided fast FFT computation. Fig. 3 plots the MMSE decoding error as a function of M for a variety of row weights L. The figure emphasizes with dashed lines the average ℓ_2 norm of x (top) and of the small coefficients (bottom); increasing M reduces the decoding error, until it reaches the energy level of the small coefficients. A small row weight, e.g., L = 5, may miss some of the large coefficients and is thus bad for decoding; as we increase L, fewer measurements are needed to obtain the same precision. However, there is an optimal $L_{\rm opt} \approx 2/S = 20$ beyond which any performance gains are marginal. Furthermore, values of $L > L_{
m opt}$ give rise to divergence in CS-BP, even with damping. An example of the output of the CS-BP decoder and how it compares to the signal x appears in Fig. 4, where we used L=20 and M=400. Although N=1000, we only plotted the first 100 signal values x(i) for ease of visualization.

To compare the performance of CS-BP with other CS decoding algorithms, we also simulated: i) ℓ_1 decoding (1) via linear programming; ii) GPSR [20], an optimization method that minimizes $||\theta||_1 + \mu||y - \Phi\Psi\theta||_2^2$; iii) CoSaMP [16], a fast greedy solver; and iv) IHT [17], an iterative thresholding algorithm. We simulated all five methods where $N=1000, S=0.1, L=20, \sigma_1=10, \sigma_0=1, p=525$, and the measurements are noiseless. Throughout the experiment we ran the different methods using the same CS-LDPC encoding matrix Φ , the same signal x, and therefore same measurements y. Fig. 5 plots the MMSE decoding error as a function of M for the five methods. For

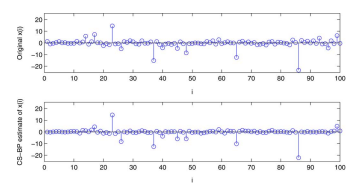


Fig. 4. Original signal x and version decoded by CS-BP. ($N=1000, S=0.1, L=20, M=400, \sigma_1=10, \sigma_0=1$, and noiseless measurements.)

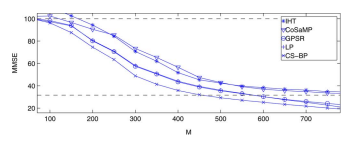


Fig. 5. MMSE as a function of the number of measurements M using CS-BP, linear programming (LP), GPSR, CoSaMP, and IHT. The dashed lines show the ℓ_2 norms of x (top) and the small coefficients (bottom). ($N=1000,S=0.1,L=20,\sigma_1=10,\sigma_0=1$, and noiseless measurements.)

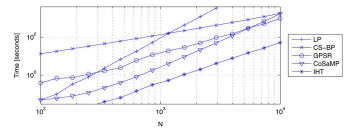


Fig. 6. Run-time in seconds as a function of the signal length N using CS-BP, linear programming (LP) ℓ_1 decoding, GPSR, CoSaMP, and IHT. ($S=0.1, L=20, M=0.4N, \sigma_1=10, \sigma_0=1$, and noiseless measurements.)

small to moderate M, CS-BP exploits its knowledge about the approximately sparse structure of x, and has a smaller decoding error. CS-BP requires 20%–30% fewer measurements than the optimization methods LP and GPSR to obtain the same MMSE decoding error; the advantage over the greedy solvers IHT and CoSaMP is even greater. However, as M increases, the advantage of CS-BP over LP and GPSR becomes less pronounced.

To compare the speed of CS-BP to other methods, we ran the same five methods as before. In this experiment, we varied the signal length N from 100 to 10 000, where $S=0.1, L=20, \sigma_1=10, \sigma_0=1, p=525$, and the measurements are noiseless. We mention in passing that some of the algorithms that were evaluated can be accelerated using linear algebra routines optimized for sparse matrices; the improvement is quite modest, and the run-times presented here do not reflect this optimization. Fig. 6 plots the run-times of the five methods in seconds as a function of N. It can be seen that LP scales more poorly than the other algorithms, and so we did not simulate it for N >

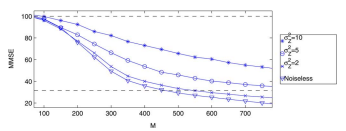


Fig. 7. MMSE as a function of M using different noise levels σ_Z^2 . The dashed lines show the ℓ_2 norms of x (top) and the small coefficients (bottom). $(N=1000,S=0.1,L=20,\sigma_1=10,$ and $\sigma_0=1.)$

3000.5 CoSaMP also seems to scale relatively poorly, although it is possible that our conjugate gradient implementation can be improved using the pseudo-inverse approach instead [16]. The run-times of CS-BP seem to scale somewhat better than IHT and GPSR. Although the asymptotic computational complexity of CS-BP is good, for signals of length N = 10000 it is still slower than IHT and GPSR; whereas IHT and GPSR essentially perform matrix-vector multiplications, CS-BP is slowed by FFT computations performed in each iteration for all nodes in the factor graph. Additionally, whereas the choice $p = O(\sigma_1/\sigma_0)$ yields $O(\sigma_1/\sigma_0 \log(\sigma_1/\sigma_0)N \log^2(N))$ complexity, FFT computation with p = 525 samples is somewhat slow. That said, our main contribution is a computationally feasible Bayesian approach, which allows to reduce the number of measurements (Fig. 5); a comparison between CS-BP and previous Bayesian approaches to CS [25], [26] would be favorable.

To demonstrate that CS-BP deals well with measurement noise, recall the noisy measurement setting $y=\Phi x+z$ of Section III, where $z\sim \mathcal{N}(0,\sigma_Z^2)$ is AWGN with variance σ_Z^2 . Our algorithm deals with noise by convolving the noiseless version of the estimated pdf (5) with the noise pdf. We simulated decoding problems where $N=1000,S=0.1,L=20,\sigma_1=10,\sigma_0=1,p=525,$ and $\sigma_Z^2\in\{0,2,5,10\}$. Fig. 7 plots the MMSE decoding error as a function of M and σ_Z^2 . To put things in perspective, the average measurement picks up a Gaussian term of variance $L(1-S)\sigma_0^2=18$ from the signal. Although the decoding error increases with σ_Z^2 , as long as $\sigma_Z^2\ll 18$ the noise has little impact on the decoding error; CS-BP offers a graceful degradation to measurement noise.

Our final experiment considers model mismatch where CS-BP has an imprecise statistical characterization of the signal. Instead of a two-state mixture Gaussian signal model as before, where large coefficients have variance σ_1^2 and occur with probability S, we defined a C-component mixture model. In our definition, σ_0^2 is interpreted as a background signal level, which appears in all coefficients. Whereas the two-state model adds a "true signal" component of variance $\sigma_1^2 - \sigma_0^2$ to the background signal, the C-1 large components each occur with probability S and the amplitudes of the true signals are $\sigma_2, 2\sigma_2, \ldots, (C-1)\sigma_2$, where σ_2 is chosen to preserve the total signal energy. At the same time, we did not change the signal priors in CS-BP, and used the same two-state mixture model as before. We simulated decoding problems where

⁵Our LP solver is based on interior point methods.

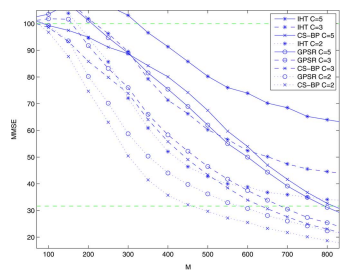


Fig. 8. MMSE as a function of the number of measurements M and the number of components C in the mixture Gaussian signal model. Plots for CS-BP (x), GPSR (circle), and IHT (asterisk) appear for C=2 (dotted), C=3 (dashed), and C=5 (solid). The horizontal dashed lines show the ℓ_2 norms of x (top) and the small coefficients (bottom). $(N=1000,S=0.1,L=20,\sigma_1=10,\sigma_0=1,$ and noiseless measurements.)

 $N=1000, S=0.1, L=20, \sigma_1=10, \sigma_0=1, p=525$, the measurements are noiseless, and $C\in\{2,3,5\}$. Fig. 8 plots the MMSE decoding error as a function of M and C. The figure also shows how IHT and GPSR perform, in order to evaluate whether they are more robust than the Bayesian approach of CS-BP. We did not simulate CoSaMP and ℓ_1 decoding, since their MMSE performance is comparable to that of IHT and GPSR. As the number of mixture components C increases, the MMSE provided by CS-BP increases. However, even for C=3 the sparsity rate effectively doubles from S to 2S, and an increase in the required number of measurements M is expected. Interestingly, the greedy IHT method also degrades significantly, perhaps because it implicitly makes an assumption regarding the number of large mixture components. GPSR, on the other hand, degrades more gracefully.

VI. VARIATIONS AND ENHANCEMENTS

Supporting Arbitrary Sparsifying Basis Ψ : Until now, we have assumed that the canonical sparsifying basis is used, i.e., $\Psi=I$. In this case, x itself is sparse. We now explain how CS-BP can be modified to support the case where x is sparse in an arbitrary basis Ψ . In the encoder, we multiply the CS-LDPC matrix Φ by Ψ^T and encode x as $y=(\Phi\Psi^T)x=(\Phi\Psi^T)(\Psi\theta)=\Phi\theta$, where $(\cdot)^T$ denotes the transpose operator. In the decoder, we use BP to form the approximation $\hat{\theta}$, and then transform via Ψ to $\hat{x}=\Psi\hat{\theta}$. In order to construct the modified encoding matrix $\Phi\Psi^T$ and later transform $\hat{\theta}$ to \hat{x} , extra computation is needed; this extra cost is $O(N^2)$ in general. Fortunately, in many practical situations Ψ is structured (e.g., Fourier or wavelet bases) and amenable to fast computation. Therefore, extending our methods to such bases is feasible.

Exploiting Statistical Dependencies: In many signal representations, the coefficients are not i.i.d. For example, wavelet representations of natural images often contain correlations be-

tween magnitudes of parent and child coefficients [2], [43]. Consequently, it is possible to decode signals from fewer measurements using an algorithm that allocates different distributions to different coefficients [46], [58]. By modifying the dependencies imposed by the prior constraint nodes (Section IV-A), CS-BP decoding supports different signal models.

Feedback: Feedback from the decoder to the encoder can be used in applications where measurements may be lost because of transmissions over faulty channels. In an analogous manner to a digital fountain [59], the marginal distributions (6) enable us to identify when sufficient information for signal decoding has been received. At that stage, the decoder notifies the encoder that decoding is complete, and the stream of measurements is stopped.

Irregular CS-LDPC Matrices: In channel coding, LDPC matrices that have irregular row and column weights come closer to the Shannon limit, because a small number of rows or columns with large weights require only modest additional computation yet greatly reduce the block error rate [38]. In an analogous manner, we expect irregular CS-LDPC matrices to enable a further reduction in the number of measurements required.

VII. DISCUSSION

This paper has developed a sparse encoding matrix and belief propagation decoding algorithm to accelerate CS encoding and decoding under the Bayesian framework. Although we focus on decoding approximately sparse signals, CS-BP can be extended to signals that are sparse in other bases, is flexible to modifications in the signal model, and can address measurement noise.

Despite the significant benefits, CS-BP is not universal in the sense that the encoding matrix and decoding methods must be modified in order to apply our framework to arbitrary bases. Nonetheless, the necessary modifications only require multiplication by the sparsifying basis Ψ or its transpose Ψ^T .

Our method resembles LDPC codes [37], [38], which use a sparse Bernoulli parity check matrix. Although any linear code can be represented as a bipartite graph, for LDPC codes the sparsity of the graph accelerates the encoding and decoding processes. LDPC codes are celebrated for achieving rates close to the Shannon limit. A similar comparison of the MMSE performance of CS-BP with information theoretic bounds on CS performance has demonstrated that CS-BP is asymptotically optimal in the large-system limit [29], [60]. Additionally, although CS-BP is not guaranteed to converge, the recent convergence proofs for LDLC codes [36] suggest that future work on extensions of CS-BP may also yield convergence proofs.

In comparison to previous work on Bayesian aspects of CS [25], [26], our method is much faster, requiring only $O(N\log^2(N))$ computation. At the same time, CS-BP offers significant flexibility, and should not be viewed as merely another fast CS decoding algorithm. However, CS-BP relies on the sparsity of CS-LDPC matrices, and future research can consider the applicability of such matrices in different applications.

APPENDIX

Outline of Proof of Theorem 1: The proof begins with a derivation of probabilistic bounds on $||x||_2$ and $||x||_{\infty}$. Next,

we review a result by Wang *et al.* [49, Theorem 1]. The proof is completed by combining the bounds with the resultby Wang *et al.*

Upper Bound on $||x||_2^2$: Consider $||x||_2^2 = \sum_{i=1}^N x_i^2$, where the random variable (RV) X_i has a mixture distribution

$$X_i^2 \sim \begin{cases} \chi^2 \sigma_1^2 & \text{w.p. } S \\ \chi^2 \sigma_0^2 & \text{w.p. } 1 - S. \end{cases}$$

Recall the moment generating function (MGF), $M_X(t) = E[e^{tx}]$. The MGF of a Chi-squared RV satisfies $M_{\chi^2}(t) = (1-2t)^{-(1)/(2)}$. For the mixture RV X_i^2 ,

$$M_{X_i^2}(t) = \frac{S}{\sqrt{1 - 2t\sigma_1^2}} + \frac{1 - S}{\sqrt{1 - 2t\sigma_0^2}}.$$

Additionally, because the X_i are i.i.d., $M_{||x||_2^2}(t) = [M_{X_i^2}(t)]^N$. Invoking the Chernoff bound, we have

$$\Pr(||x||_{2}^{2} < SN\sigma_{1}^{2}) < e^{-tSN\sigma_{1}^{2}} \left[\frac{S}{\sqrt{1 - 2t\sigma_{1}^{2}}} + \frac{1 - S}{\sqrt{1 - 2t\sigma_{0}^{2}}} \right]^{N}$$

for t<0. We aim to show that $\Pr(||x||_2^2 < SN\sigma_1^2)$ decays faster than $N^{-\gamma}$ as N is increased. To do so, let $t=-\alpha/(\sigma_1^2)$, where $\alpha>0$. It suffices to prove that there exists some α for which

$$f_1(\alpha) = e^{\alpha S} \left[\frac{S}{\sqrt{1+2\alpha}} + \frac{1-S}{\sqrt{1+2\alpha \left(\frac{\sigma_0}{\sigma_1}\right)^2}} \right] < 1.$$

Let $f_2(\alpha) = 1/\sqrt{1+2\alpha}$ and $f_3(\alpha) = e^{\alpha}$. It is easily seen via Taylor series that $f_2(\alpha) = 1 - \alpha + O(\alpha^2)$ and $f_3(\alpha) = 1 + \alpha + O(\alpha^2)$, and so

$$f_{1}(\alpha) = e^{\alpha S} \left[S(1 - \alpha + O(\alpha^{2})) + (1 - S) \left(1 - \alpha \left(\frac{\sigma_{0}}{\sigma_{1}} \right)^{2} + O\left(\alpha^{2} \left(\frac{\sigma_{0}}{\sigma_{1}} \right)^{4} \right) \right) \right]$$

$$= \left[1 + \alpha S + O(\alpha^{2} S^{2}) \right] \left[1 - \alpha \left(S + (1 - S) \left(\frac{\sigma_{0}}{\sigma_{1}} \right)^{2} \right) + O(\alpha^{2}) \right].$$

Because of the negative term $-\alpha(1-S)(\sigma_0/\sigma_1)^2 < 0$, which dominates the higher order term $O(\alpha^2)$ for small α , there exists $\alpha>0$, which is independent of N, for which $f_1(\alpha)<1$. Using this α , the Chernoff bound provides an upper bound on $\Pr(||x||_2^2 < SN\sigma_1^2)$ that decays exponentially with N. In summary,

$$\Pr(||x||_2^2 < SN\sigma_1^2) = o(N^{-\gamma}).$$
 (8)

Lower Bound on $||x||_2^2$: In a similar manner, MGF's and the Chernoff bound can be used to offer a probabilistic bound on the number of large Gaussian mixture components

$$\Pr\left(\sum_{i=1}^{N} Q(i) > \frac{3}{2}SN\right) = o(N^{-\gamma}). \tag{9}$$

Taking into account the limited number of large components and the expected squared ℓ_2 norm, $E[||x||_2^2] = N[S\sigma_1^2 + (1-S)\sigma_0^2]$, we have

$$\Pr(||x||_2^2 > N[2S\sigma_1^2 + (1-S)\sigma_0^2]) = o(N^{-\gamma}).$$
 (10)

We omit the (similar) details for brevity.

Bound on $||x||_{\infty}$: The upper bound on $||x||_{\infty}$ is obtained by first considering large mixture components and then small components. First, we consider the large Gaussian mixture components, and denote $x_L = \{x(i) : Q(i) = 1\}$.

$$\Pr\left(\|x_L\|_{\infty} < \sqrt{2\ln(SN^{1+\gamma})}\sigma_1 \left| \sum_{i=1}^{N} Q(i) \le \frac{3}{2}SN \right.\right)$$

$$\ge \left[f_4(\sqrt{2\ln(SN^{1+\gamma})})\right]^{\frac{3}{2}SN} \tag{11}$$

$$> \left[1 - \frac{f_5\left(\sqrt{2\ln(SN^{1+\gamma})}\right)}{\sqrt{2\ln(SN^{1+\gamma})}}\right]^{\frac{3}{2}SN} \tag{12}$$

$$> 1 - \frac{3}{2}SN \frac{f_5\left(\sqrt{2\ln(SN^{1+\gamma})}\right)}{\sqrt{2\ln(SN^{1+\gamma})}} \tag{13}$$

$$= 1 - \frac{3SN}{2\sqrt{2\ln(SN^{1+\gamma})}} \frac{e^{-\frac{1}{2}2\ln(SN^{1+\gamma})}}{\sqrt{2\pi}}$$

$$= 1 - \frac{3N^{-\gamma}}{4\sqrt{\ln(SN^{1+\gamma})}}$$

where $f_4(\alpha)=(2\pi)^{-1/2}\int_{-\infty}^{\alpha}e^{-u^2/2}du$ is the cumulative distribution function of the standard normal distribution, the inequality (11) relies on $f_4(\cdot)<1$ and the possibility that $\sum_{i=1}^N Q(i)$ is strictly smaller than $(3)/(2)SN, f_5(\alpha)=(1)/(\sqrt{2\pi})e^{-\alpha^2/2}$ is the pdf of the standard normal distribution, (12) relies on the bound $f_4(\alpha)>1-f_5(\alpha)/\alpha$, and the inequality (13) is motivated by $(1-\alpha)^\beta>1-\alpha\beta$ for $\alpha,\beta>0$. Noting that $\ln(SN^{1+\gamma})$ increases with N, for large N we have

$$\Pr\left(\|x_L\|_{\infty} < \sqrt{2\ln(SN^{1+\gamma})}\sigma_1 \left| \sum_{i=1}^N Q(i) \le \frac{3}{2}SN \right.\right)$$
$$> 1 - \frac{N^{-\gamma}}{5}. \quad (14)$$

Now consider the small Gaussian mixture components, and denote $x_S = \{x(i) : Q(i) = 0\}$. As before

$$\Pr\left(||x_S||_{\infty} < \sqrt{2\ln(SN^{1+\gamma})}\sigma_1\right)$$

$$\geq \left[f_4\left(\sqrt{2\ln(SN^{1+\gamma})}\frac{\sigma_1}{\sigma_0}\right)\right]^N$$

$$> 1 - \frac{N}{\sqrt{2\ln(SN^{1+\gamma})}\frac{\sigma_1}{\sigma_0}} \frac{e^{-\frac{1}{2}2\ln(SN^{1+\gamma})\left(\frac{\sigma_1}{\sigma_0}\right)^2}}{\sqrt{2\pi}} \quad (15)$$

where in (15) the number of small mixture components is often less than N. Because $\sigma_1 > \sigma_0$, for large N we have

$$\Pr(||x_S||_{\infty} < \sqrt{2\ln(SN^{1+\gamma})}\sigma_1) > 1 - \frac{N^{-\gamma}}{5}.$$
 (16)

Combining (9), (14) and (16), for large N we have

$$\Pr\left(||x||_{\infty} < \sqrt{2\ln(SN^{1+\gamma})}\sigma_1\right) > 1 - \frac{N^{-\gamma}}{2}.\tag{17}$$

Result by Wang et al. ([49], Theorem 1):

Theorem 2—[49]: Consider $x \in \mathbb{R}^N$ that satisfies the condition

$$\frac{\|x\|_{\infty}}{\|x\|_2} \le Q. \tag{18}$$

In addition, let V be any set of N vectors $\{v_1, \ldots, v_N\} \subset \mathbb{R}^N$. Suppose a sparse random matrix $\Phi \in \mathbb{R}^{M \times N}$ satisfies

$$E[\Phi_{ij}] = 0, E\left[\Phi_{ij}^2\right] = 1, E\left[\Phi_{ij}^4\right] = s,$$

where (1)/(s) = (L)/(N) is the fraction of nonzero entries in Φ . Let

$$M = \begin{cases} O\left(\frac{1+\gamma}{\epsilon^2} sQ^2 \log(N)\right) & \text{if } sQ^2 \ge \Omega(1) \\ O\left(\frac{1+\gamma}{\epsilon^2} \log(N)\right) & \text{if } sQ^2 \le O(1). \end{cases}$$
(19)

Then with probability at least $1 - N^{-\gamma}$, the random projections $\Phi x/M$ and $\Phi v_i/M$ can produce an estimate \hat{a}_i for $x^T v_i$ satisfying

$$|\hat{a}_i - x^T v_i| \le \epsilon ||x||_2 ||v_i||_2, \quad \forall i \in \{1, \dots, N\}.$$

Application of Theorem 2 to Proof of Theorem 1: Combining (8), (10), and (17), the union bound demonstrates that with probability lower bounded by $1-N^{-\gamma}$ we have $||x||_{\infty} < \sqrt{2\ln(SN^{1+\gamma})}\sigma_1$ and $||x||_2^2 \in (NS\sigma_1^2,N[2S\sigma_1^2+(1-S)\sigma_0^2]).6$ When these ℓ_2 and ℓ_∞ bounds hold, we can apply Theorem 2.

To apply Theorem 2, we must specify (i) Q (18); ii) the test vectors $(v_i)_{i=1}^N$; iii) the matrix sparsity s; and iv) the ϵ parameter. First, the bounds on $\|x\|_2$ and $\|x\|_\infty$ indicate that $\|x\|_\infty/\|x\|_2 \leq Q = \sqrt{(2\ln(SN^{1+\gamma}))/(SN)}$. Second, we choose $(v_i)_{i=1}^N$ to be the N canonical vectors of the identity matrix I_N , providing $x^Tv_i = x_i$. Third, our choice of L offers $s = N/L = NS/(\eta \ln(SN^{1+\gamma}))$. Fourth, we set

 $^6 {\rm The} \ o(\,\cdot\,)$ terms (8) and (10) demonstrate that there exists some N_0 such that for all $N>N_0$ the upper and lower bounds on $\|x\|_2^2$ each hold with probability lower bounded by $1-(1)/(4)N^\gamma$, resulting in a probability lower bounded by $1-N^{-\gamma}$ via the union bound. Because the expression (2) for the number of measurements M is an order term, the case where $N\leq N_0$ is inconsequential.

$$\epsilon = \frac{\mu \sigma_1}{\sqrt{N[2S\sigma_1^2 + (1-S)\sigma_0^2]}}$$

Using these parameters, Theorem 2 demonstrates that all N approximations \hat{a}_i satisfy

$$|\hat{a}_i - x_i| = |\hat{a}_i - x^T v_i| \le \epsilon ||x||_2 ||v_i||_2 < \mu \sigma_1$$

with probability lower bounded by $1 - N^{-\gamma}$. Combining the probability that the ℓ_2 and ℓ_{∞} bounds hold and the decoding probability offered by Theorem 2, we have

$$\|\hat{a} - x\|_{\infty} < \mu \sigma_1 \tag{20}$$

with probability lower bounded by $1 - 2N^{-\gamma}$.

We complete the proof by computing the number of measurements ${\cal M}$ required (19). Because

$$sQ^2 = \frac{K}{\eta \ln(SN^{1+\gamma})} \frac{2\ln(SN^{1+\gamma})}{SN} = \frac{2}{\eta}$$

we need

$$M = O\left((1 + 2\eta^{-1})\frac{1 + \gamma}{\epsilon^2}\log(N)\right)$$

$$= O\left(N(1 + 2\eta^{-1})\frac{(1 + \gamma)}{\mu^2}\left[2S + (1 - S)\left(\frac{\sigma_0}{\sigma_1}\right)^2\right]\log(N)\right)$$
measurements.

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