

BlockTensorDecompositions.jl: A Unified Constrained Tensor Decomposition Julia Package

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1 Introduction

- Tenors are useful in many applications
- Need tools for fast and efficient decompositions

For the scientific user, it would be most useful for there to be a single piece of software that can take as input 1) any reasonable type of factorization model and 2) constraints on the individual factors, and produce a factorization. Details like what rank to select, how the constraints should be enforced, and convergence criteria should be handled automatically, but customizable to the knowledgeable user. These are the core specification for BlockTensorDecompositions.jl.

1.1 Related tools

- Packages within Julia
- Other languages
- Hint at why I developed this

Beyond the external usefulness already mentioned, this package offers a playground for fair comparisons of different parameters and options for performing tensor factorizations across various decomposition models. There exist packages for working with tensors in languages like Python (TensorFlow [1], PyTorch [2], and TensorLy [3]), MATLAB (Tensor Toolbox [4]), R (rTensor [5]), and Julia (TensorKit.jl [6], Tullio.jl [7], OMEinsum.jl [8], and TensorDecompositions.jl [9]). But they only provide a groundwork for basic manipulation of tensors and the most common tensor decomposition models and algorithms, and are not equipped to handle arbitrary user defined constraints and factorization models.

Some progress towards building a unified framework has been made [10–12]. But these approaches don't operate on the high dimensional tensor data natively and rely on matricizations of the problem, or only consider nonnegative constraints. They also don't provide an all-in-one package for executing their frameworks.

1.2 Contributions

- Fast and flexible tensor decomposition package
- Framework for creating and performing custom
 - tensor decompositions
 - constrained factorization (the what)
 - iterative updates (the how)
- Implement new “tricks”
 - a (Lipschitz) matrix stepsize for efficient sub-block updates
 - multi-scaled factorization when tensor entries are discretizations of a continuous function
 - partial projection and rescaling to enforce linear constraints (rather than Euclidean projection)
- ?? rank detection ??

The main contribution is a description of a fast and flexible tensor decomposition package, along with a public implementation written in Julia: BlockTensorDecompositions.jl. This package provides a framework for creating and performing custom tensor decompositions. To the author’s knowledge, it is the first package to provide automatic factorization to a large class of constrained tensor decompositions problems, as well as a framework for implementing new constraints and iterative algorithms. This paper also describes three new techniques not found in the literature that empirically converge faster than traditional block-coordinate descent.

2 Tensor Decompositions

- the math section of the paper

This section reviews the notation used throughout the paper and commonly used tensor decompositions.

2.1 Notation

- tensor notation, use MATLAB notation for indexing so subscripts can be used for a sequence of tensors

2.1.a Sets

The set of real numbers is denoted as \mathbb{R} and its restrictions to nonnegative numbers is denoted as $\mathbb{R}_+ = \mathbb{R}_{\geq 0} = \{x \in \mathbb{R} \mid x \geq 0\}$.

We use $[N] = \{1, 2, \dots, N\} = \{n\}_{n=1}^N$ to denote integers from 1 to N .

Usually, lower case symbols will be used for the running index, and the capitalized letter will be the maximum letter it runs to. This leads to the convenient shorthand $i \in [I]$, $j \in [J]$, etc.

We use a capital delta Δ to denote sets of vectors or higher order tensors where the slices or fibres along a specified dimension sum to 1, i.e. generalized simplexes.

Usually, we use script letters ($\mathcal{A}, \mathcal{B}, \mathcal{C}$, etc.) for other sets.

2.1.b Vectors, Matrices, and Tensors

Vectors are denoted with lowercase letters (x, y , etc.), and matrices and higher order tensors with uppercase letters (commonly A, B, C and X, Y, Z). The order of a tensor is the number of axes it

has. We would call vectors “order-1” or “1st order” tensors, and matrices “order-2” or “2nd order” tensors.

To avoid confusion between entries of a vector/matrix/tensor and indexing a list of objects, we use square brackets to denote the former, and subscripts to denote the later. For example, the entry in the i th row and j th column of a matrix $A \in \mathbb{R}$ is $A[i, j]$. This follows MATLAB/Julia notation where $A[i, j]$ points to the entry $A[i, j]$. We contrast this with a list of I objects being denoted as a_1, \dots, a_I , or more compactly, $\{a_i\}$ when it is clear the index $i \in [I]$.

The transpose $A^\top \in \mathbb{R}^{J \times I}$ of a matrix $A \in \mathbb{R}^{I \times J}$ flips entries along the main diagonal: $A^\top[j, i] = A[i, j]$. In Julia, the transpose of a matrix is typed with a single apostrophe A' .

The n -slices, n th mode slices, or mode n slices of an N th order tensor A are notated with the slice $A[:, \dots, :, i_n, :, \dots, :]$. For a 3rd order tensor A , the 1st, 2nd, and 3rd mode slices $A[i, :, :]$, $A[:, j, :]$, and $A[:, :, k]$ have special names and are called the horizontal, lateral, and frontal slices and are displayed in Figure 1. In Julia, the 1-, 2-, and 3-slices of a third order array A would be `eachslice(A, dims=1)`, `eachslice(A, dims=2)`, and `eachslice(A, dims=3)`.

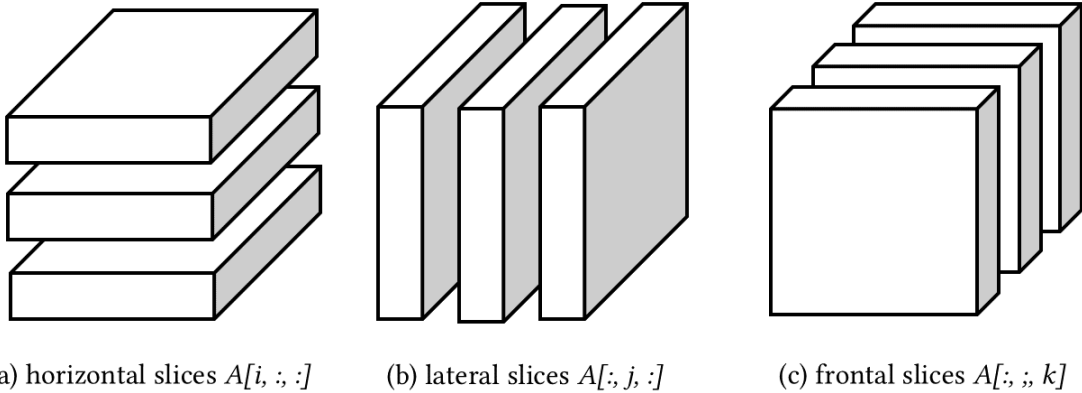


Figure 1: Slices of an order 3 tensor A .

The n -fibres, n th mode fibres, or mode n fibres of an N th order tensor A are denoted $A[i_1, \dots, i_{n-1}, :, i_{n+1}, \dots, i_N]$. For example, the 1-fibres of a matrix M are the column vectors $M[:, j]$, and the 2-fibres are the row vectors $M[i, :]$. For order-3 tensors, the 1st, 2nd, and 3rd mode fibres $A[:, j, k]$, $A[i, :, :]$, and $A[i, j, :]$ are called the vertical/column, horizontal/row, and depth/tube fibres respectively and are displayed in Figure 2. Natively in Julia, the 1-, 2-, and 3-fibres of a third order array A would be `eachslice(A, dims=(2,3))`, `eachslice(A, dims=(1,3))`, and `eachslice(A, dims=(1,2))`. BlockTensorDecomposition.jl defines the function `eachfibre(A; n)` to do exactly this. For example, the 1-fibres of an array A would be `eachfibre(A, n=1)`.

For matrices, the 1-fibres are the same as the 2-slices (and vice versa), but for N th order tensors in general, fibres are always vectors, whereas n -slices are $(N - 1)$ th order tensors.

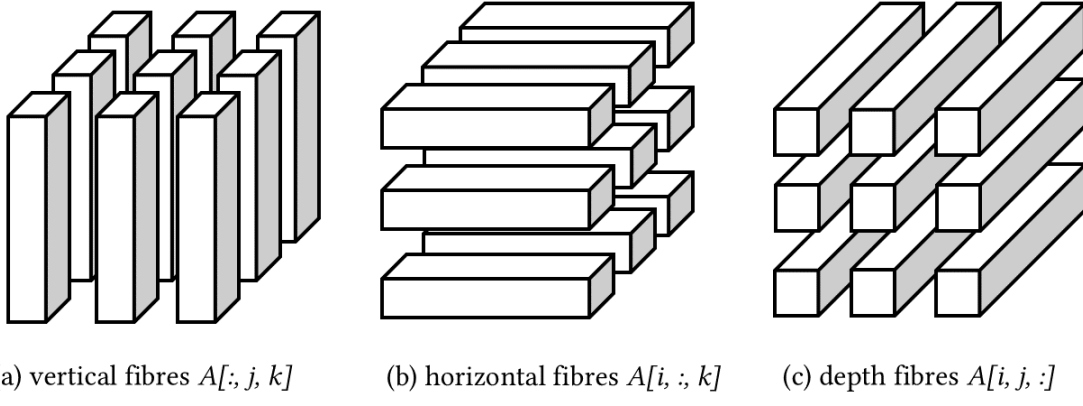


Figure 2: Fibres of an order 3 tensor A .

Since we commonly use I as the size of a tensor's dimension, we use id_I to denote the identity tensor of size I (of the appropriate order). When the order is 2, id_I is an $I \times I$ matrix with ones along the main diagonal, and zeros elsewhere. For higher orders N , this is an $\underbrace{I \times \cdots \times I}_{N \text{ times}}$ tensor where $\text{id}_I[i_1, \dots, i_N] = 1$ when $i_1 = \dots = i_N \in [I]$, and is zero otherwise.

`BlockTensorDecomposition.jl` defines `identity_tensor(I, ndims)` to construct id_I .

For a vector, matrix, or tensor filled with ones, we use $\mathbb{1} \in \mathbb{R}^{I_1 \times \cdots \times I_N}$. This can be constructed in Julia with `ones(I1, ..., In)`.

2.1.c Products of Tensors

Definition 2.1: The outer product \otimes between two tensors $A \in \mathbb{R}^{I_1 \times \cdots \times I_M}$ and $B \in \mathbb{R}^{J_1 \times \cdots \times J_N}$ yields an order $M + N$ tensor $A \otimes B \in \mathbb{R}^{I_1 \times \cdots \times I_M \times J_1 \times \cdots \times J_N}$ that is entry-wise

$$(A \otimes B)[i_1, \dots, i_M, j_1, \dots, j_N] = A[i_1, \dots, i_M]B[j_1, \dots, j_N].$$

TODO Define in `BlockTensorDecomposition.jl`

The Frobenius inner product between two tensors $A, B \in \mathbb{R}^{I_1 \times \cdots \times I_N}$ yields a real number $A \cdot B \in \mathbb{R}$ and is defined as

$$\langle A, B \rangle = A \cdot B = \sum_{i_1=1}^{I_1} \cdots \sum_{i_N=1}^{I_N} A[i_1, \dots, i_N]B[i_1, \dots, i_N].$$

Julia's standard library package `LinearAlgebra` implements the Frobenius inner product with `dot(A, B)` or `A' * B`.

The n -slice dot product \cdot_n between two tensors $A \in \mathbb{R}^{K_1, \dots, K_{n-1}, I, K_{n+1}, \dots, K_N}$ and $B \in \mathbb{R}^{K_1, \dots, K_{n-1}, J, K_{n+1}, \dots, K_N}$ returns a matrix $(A \cdot_n B) \in \mathbb{R}^{I \times J}$ with entries

$$(A \cdot_n B)[i, j] = \sum_{k_1 \dots k_{n-1} k_{n+1} \dots k_N} A[k_1, \dots, k_{n-1}, i, k_{n+1}, \dots, k_N] B[k_1, \dots, k_{n-1}, j, k_{n+1}, \dots, k_N].$$

This product can also be thought of as taking the dot product $(A \cdot_n B)[i, j] = A_i \cdot B_j$ between all pairs of n th order slices of A and B , which exactly how BlockTensorDecomposition.jl defines the operation.

```

function slicewise_dot(A::AbstractArray, B::AbstractArray; dims=1)
    C = zeros(size(A, dims), size(B, dims))
    if A === B # use faster routine if they are the same
        return _slicewise_self_dot!(C, A; dims)
    end

    for (i, A_slice) in enumerate(eachslice(A; dims))
        for (j, B_slice) in enumerate(eachslice(B; dims))
            C[i, j] = A_slice ∙ B_slice
        end
    end
    return C
end

function _slicewise_self_dot!(C, A; dims=1)
    enumerated_A_slices = enumerate(eachslice(A; dims))
    for (i, Ai_slice) in enumerated_A_slices
        for (j, Aj_slice) in enumerated_A_slices
            if i > j
                continue
            else # only compute the upper triangle entries of C
                C[i, j] = Ai_slice ∙ Aj_slice
            end
        end
    end
    return Symmetric(C) # indexing C[2,1] points to the entry in C[1,2]
end

```

BlockTensorDecomposition.jl defines this operation with `slicewise_dot(A, B, n)`. In the special case where $A = B$, a more efficient method that only computes entries where $i \leq j$ is defined since $A \cdot_n A$ is a symmetric matrix.

The n -slice product of a tensor with itself $X \cdot_n X$ should be thought of as a generalization of the Gram matrix $X^\top X$ since it considers the matrix generated by taking the dot product between every n th mode slice, just like how the Gram matrix considers the dot product between every pair of columns.

The n -mode product \times_n between a tensor $A \in \mathbb{R}^{I_1 \times \dots \times I_N}$ and matrix $B \in \mathbb{R}^{I_n \times J}$, returns a tensor $(A \times_n B) \in \mathbb{R}^{I_1 \times \dots \times I_{n-1} \times J \times I_{n+1} \times \dots \times I_N}$ with entries

$$(A \times_n B)[i_1, \dots, i_{n-1}, j, i_{n+1}, \dots, i_N] = \sum_{i_n=1}^{I_n} A[i_1, \dots, i_{n-1}, i_n, i_{n+1}, \dots, i_N] B[i_n, j].$$

BlockTensorDecomposition.jl defines this operation with `nmode_product(A, B, n)`.

```

function nmode_product(A::AbstractArray, B::AbstractMatrix, n::Integer)
    # convert the problem to the mode-1 product
    Aperm = swapdims(A, n)
    Cperm = Aperm ×₁ B
    return swapdims(Cperm, n) # swap back
end

function ×₁(А::AbstractArray, B::AbstractMatrix)
    # Turn the 1-mode product into matrix-matrix multiplication
    sizeA = size(A)
    Amat = reshape(A, sizeA[1], :)

    # Initialize the output tensor
    C = zeros(size(B, 1), sizeA[2:end]...)
    Cmat = reshape(C, size(B, 1), prod(sizeA[2:end])))

    # Perform matrix-matrix multiplication Cmat = B*Amat
    mul!(Cmat, B, Amat)

    return C # Output entries of Cmat in tensor form
end

function swapdims(A::AbstractArray, a::Integer, b::Integer=1)
    # Construct a permutation where a and b are swapped
    # e.g. [4, 2, 3, 1, 5, 6] when a=4 and b=1
    dims = collect(1:ndims(A))
    dims[a] = b; dims[b] = a
    return permutedims(A, dims)
end

```

! Note

If we were only working with a fixed order of tensors, we could have defined \times_1 entry-wise with Tullio.jl. The function definition `tullio×1` below gives an example for order three tensors.

```
function tullio×1(A::AbstractArray{_,3}, B::AbstractMatrix)
    @tullio C[i, j, k] := A[r, j, k] * B[i, r]
    return C
end
```

But we would need a new definition for each ordered tensor, or use Julia's meta programming to write a method for each order at runtime.

The n -mode product and n -slice product can be thought of as opposites of each other. The n -mode product sums over just the n th dimension of the first tensor, whereas the n -slice product sums over all but the n th dimension.

We can extend the n -mode product to sum over multiple indices between two tensors.

The multi-mode product $\times_{1,\dots,n} = \times_{1:n} = \times_{[n]}$ between a tensor $A \in \mathbb{R}^{I_1 \times \dots \times I_N}$ and tensor $B \in \mathbb{R}^{I_1 \times \dots \times I_n}$, returns a tensor $(A \times_{[n]} B) \in \mathbb{R}^{I_{n+1} \times \dots \times I_N}$ with entries

$$(A \times_{[n]} B)[i_{n+1}, \dots, i_N] = \sum_{i_1, \dots, i_n} A[i_1, \dots, i_n, i_{n+1}, \dots, i_N] B[i_1, \dots, i_n].$$

This product contracts the first n indexes of A with every index of B .

More generally, we can contract any number of indexes such as the last n indexes of A with every index of B with $\times_{N-n+1, \dots, N} = \times_{(N-n+1):N} = \times_{[-n]}$,

$$(A \times_{[-n]} B)[i_1, \dots, i_n] = \sum_{i_{n+1}, \dots, i_N} A[i_1, \dots, i_{N-n}, i_{N-n+1}, \dots, i_N] B[i_{N-n+1}, \dots, i_N],$$

or specific indexes. For example, we would define $(A \times_{1,3,5} B) \in \mathbb{R}^{I_2 \times I_4 \times I_6}$ where $A \in \mathbb{R}^{I_1 \times \dots \times I_6}$ and $B \in \mathbb{R}^{I_1 \times I_3 \times I_5}$ to be

$$(A \times_{1,3,5} B)[i_2, i_4, i_6] = \sum_{i_1, i_3, i_5} A[i_1, i_2, i_3, i_4, i_5, i_6] B[i_1, i_3, i_5].$$

When A a *half-symmetric*¹ tensor of order $2N$

$$A[i_1, \dots, i_N, i_{N+1}, \dots, i_{2N}] = A[i_{N+1}, \dots, i_{2N}, i_1, \dots, i_N], \quad (1)$$

we have

¹For example, the Hessian of a scalar function (see Definition 2.3).

$$A \times_{[N]} B = A \times_{[-N]} B$$

for tensors B of order N .

2.1.d Gradients, Norms, and Lipschitz Constants

Definition 2.2 (Gradient): The gradient $\nabla f : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}^{I_1 \times \dots \times I_N}$ of a (differentiable) function $f : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}$ is defined entry-wise for a tensor $A \in \mathbb{R}^{I_1 \times \dots \times I_N}$ by

$$\nabla f(A)[i_1, \dots, i_N] = \frac{\partial f}{\partial A[i_1, \dots, i_N]}(A).$$

Definition 2.3 (Hessian): The Hessian $\nabla^2 f : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}^{(I_1 \times \dots \times I_N)^2}$ of a second-differentiable function $f : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}$ is the gradient of the gradient and is defined for a tensor $A \in \mathbb{R}^{I_1 \times \dots \times I_N}$ entry-wise by

$$\nabla^2 f(A)[i_1, \dots, i_N, j_1, \dots, j_N] = \frac{\partial^2 f}{\partial A[i_1, \dots, i_N] \partial A[j_1, \dots, j_N]}(A).$$

For a tensor input A of order N , the Hessian tensor $\nabla^2 f(A)$ is of order $2N$.

See Section 8.1 for how this definition can be reproduced by performing two gradients $\nabla^2 f = \nabla(\nabla f)$.

Definition 2.4: The Frobenius norm of a tensor A is the square root of its dot product with itself

$$\|A\|_F = \sqrt{\langle A, A \rangle}.$$

For vectors v , this is equivalent to the (Euclidean) 2-norm

$$\|v\|_F = \|v\|_2 = \sqrt{\langle v, v \rangle}.$$

For matrices M , the $(2 \rightarrow 2)$ operator norm is defined as

$$\|M\|_{\text{op}} = \sup_{\|v\|_2=1} \|Mv\|_2 = \sigma_1(M)$$

where $\sigma_1(M)$ is the largest singular value of M .

For tensors T , the operator-norm is ambiguous since there are multiple ways we can treat tensors as function on other tensors. There is a canonical way to do this for vectors $x \mapsto v^\top x$ and matrices

$x \mapsto Mx$, but not tensors. In the case of the Hessian tensor $\nabla^2 f(A) \in \mathbb{R}^{(I_1 \times \cdots \times I_N)^2}$ evaluated at $A \in \mathbb{R}^{I_1 \times \cdots \times I_N}$, it is natural to consider the function $X \mapsto \nabla^2 f(A) \times_{[N]} X$ for $X \in \mathbb{R}^{I_1 \times \cdots \times I_N}$. This gives us our definition of the operator norm on tensors.

Definition 2.5 (Operator Norm): The operator norm of a half-symmetric tensor $A \in \mathbb{R}^{(I_1 \times \cdots \times I_N)^2}$ (Equation 1) is defined as

$$\|A\|_{\text{op}} = \sup_{\|X\|_F=1} \|A \times_{[N]} X\|_F. \quad (2)$$

In Equation 2, $X \in \mathbb{R}^{I_1 \times \cdots \times I_N}$. Note that this definition agrees with the usual operator norm on matrices when $N = 1$.

Theorem 2.1 (Norm of an outer product): Let $T = A_1 \otimes \cdots \otimes A_N \in \mathbb{R}^{(I_1 \times \cdots \times I_N)^2}$ where $A_n \in \mathbb{R}^{I_n \times I_n}$ are symmetric matrices.

Then T is half-symmetric and

$$\|T\|_{\text{op}} = \prod_{n=1}^N \|A_n\|_{\text{op}}.$$

⚠️ Warning

According to how the outer product \otimes is defined in Definition 2.1, the product $A_1 \otimes \cdots \otimes A_N$ shown in Theorem 2.1 is really an element of $\mathbb{R}^{I_1 \times I_1 \times \cdots \times I_N \times I_N}$. Note how the indexes are ordered differently than an element of $\mathbb{R}^{(I_1 \times \cdots \times I_N)^2} = \mathbb{R}^{I_1 \times \cdots \times I_N \times I_1 \times \cdots \times I_N}$. Correcting for this with explicit notation becomes cumbersome and would require tensor transposes, a new definition of an outer product, or reordering of indexes in the definition of a half-symmetric tensor. These can have knock-on effects to the definition of the Hessian, multi-mode product, and the operator norm.

To avoid the headache, the equality

$$T = A_1 \otimes \cdots \otimes A_N$$

in Theorem 2.1 should be thought of as the following entry-wise equation

$$T[i_1, \dots, i_N, j_1, \dots, j_N] = A_1[i_1, j_1] \cdots A_N[i_N, j_N]. \quad (3)$$

With the outer product understood as Equation 3, the results of Theorem 2.1 that T is half-symmetric and its operator norm is the product of the operator norms of the constituent matrices is true.

Definition 2.6 (Lipschitz Function): A function $g : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}^{I_1 \times \dots \times I_N}$ is L -Lipschitz when

$$\|g(A) - g(B)\|_F \leq L\|A - B\|_F, \quad \forall A, B \in \mathbb{R}^{I_1 \times \dots \times I_N}.$$

We call the smallest such L the Lipschitz constant of g .

Definition 2.7 (Smooth Function): A differentiable function $f : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}$ is L -smooth when its gradient $g = \nabla f$ is L -Lipschitz.

Theorem 2.2 (Quadratic Smoothness): Let $f : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}$ be a quadratic function

$$f(X) = \frac{1}{2}A(X, X) + B(X) + C$$

of X with bilinear function $A : (\mathbb{R}^{I_1 \times \dots \times I_N})^2 \rightarrow \mathbb{R}$, linear function $B : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}$, and constant $C \in \mathbb{R}^{I_1 \times \dots \times I_N}$.

Then:

1. the Hessian $\nabla^2 f$ is a constant function that evaluates to $\nabla^2 f(X) = D$ at every point X for some $D \in \mathbb{R}^{(I_1 \times \dots \times I_N)^2}$,
2. the tensor D only depends on the bilinear function A (and not on B and C), and
3. the quadratic function f is L -smooth with constant

$$L = \|D\|_{\text{op}}.$$

2.2 Common Decompositions

- Extensions of PCA/ICA/NMF to higher dimensions
- talk about the most popular Tucker, Tucker-n, CP
- other decompositions
 - high order SVD (see Kolda and Bader)
 - HOSVD (see Kolda, Shifted power method for computing tensor eigenpairs)

A tensor decomposition is a factorization of a tensor into multiple (usually smaller) tensors, that can be recombined into the original tensor. To make a common interface for decompositions, we make an abstract subtype of Julia's `AbstractArray`, and subtype `AbstractDecomposition` for our concrete tensor decompositions.

```
abstract type AbstractDecomposition{T, N} <: AbstractArray{T, N} end
```

Computationally, we can think of a generic decomposition as storing factors (A, B, C, \dots) and operations ($\times_a, \times_b, \dots$) for combining them. This is what we do in BlockTensorDecomposition.jl.

```
struct GenericDecomposition{T, N} <: AbstractDecomposition{T, N}
    factors::Tuple{Vararg{AbstractArray{T}}}
    contractions::Tuple{Vararg{Function}}
end
# Y = A ×₁ B ×₂ C
array(G::GenericDecomposition) = multifoldl(contractions(G), factors(G))
```

The function `multifoldl` applies the given operations between each factor, from left to right.

```
function multifoldl(ops, args)
    @assert (length(ops) + 1) == length(args)
    x, xs... = args
    for (op, arg) in zip(ops, xs)
        x = op(x, arg)
    end
    return x
end
```

Different types of decompositions define different operations, and different “ranks” of the same decomposition specific the sizes of the factors used.

A commonly used family of decompositions can be derived from the Tucker decomposition.

Definition 2.8: A rank- (R_1, \dots, R_N) Tucker decomposition of a tensor $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$ produces N matrices $A_n \in \mathbb{R}^{I_n \times R_n}$, $n \in [N]$, and core tensor $B \in \mathbb{R}^{R_1 \times \dots \times R_N}$ such that

$$Y[i_1, \dots, i_N] = \sum_{r_1=1}^{R_1} \dots \sum_{r_N=1}^{R_N} A_1[i_1, r_1] \cdots A_r[i_N, r_N] B[r_1, \dots, r_N] \quad (4)$$

entry-wise. More compactly, this decomposition can be written using the n -mode product, or with double brackets

$$Y = B \times_1 A_1 \times_2 \dots \times_N A_N = B \bigtimes_n A_n = \llbracket B; A_1, \dots, A_N \rrbracket. \quad (5)$$

The *Tucker Product* defined by Equation 5 is implemented in BlockTensorDecomposition.jl with `tuckerproduct(B, (A1, ..., AN))` and computes

$$B \bigtimes_n A_n = \llbracket B; A_1, \dots, A_N \rrbracket.$$

It can also optionally “exclude” one of the matrix factors with the call `tuckerproduct(B, (A1, ..., AN); exclude=n)` to compute

$$B \bigtimes_{m \neq n} A_m = [B; A_1, \dots, A_{n-1}, \text{id}_{R_n}, A_{n+1}, \dots, A_N].$$

```
function tuckerproduct(core, matrices; exclude=nothing)
    N = ndims(core)
    if isnothing(exclude)
        return multifoldl(tucker_contractions(N), (core, matrices...))
    else
        return multifoldl(getnotindex(tucker_contractions(N), exclude), (core,
getnotindex(matrices, exclude)...))
    end
end

tucker_contractions(N) = Tuple((B, A) -> nmode_product(B, A, n) for n in 1:N)
```

Sometimes we write $A_0 = B$ to ease notation, and suggest the “zeroth” factor of the tucker decomposition is the core tensor B . In the special case when $N = 3$, we can visualize Tucker decomposition as multiplying the core tensor by matrices on all three sides as shown in Figure 3.

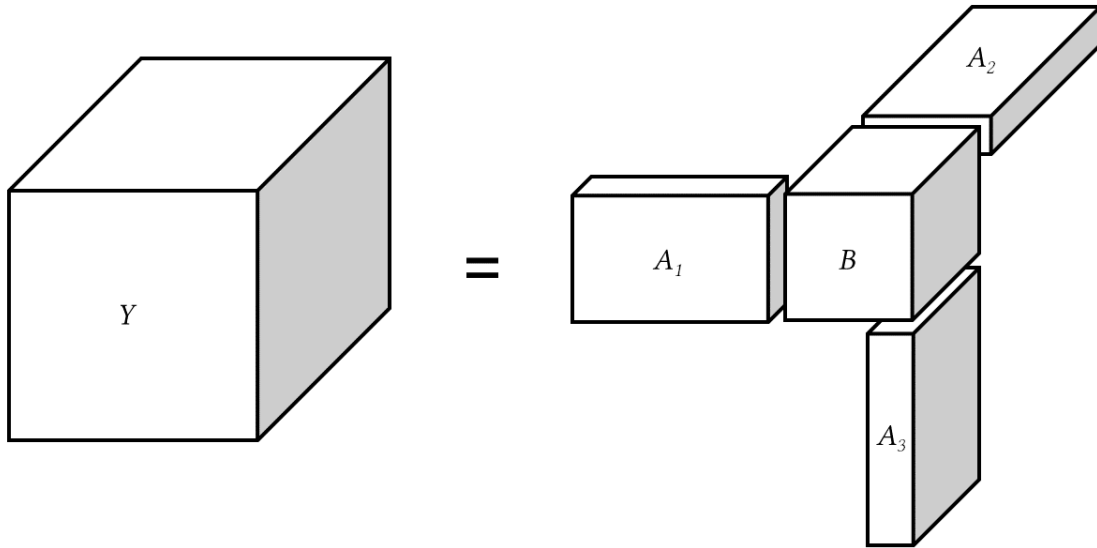


Figure 3: Tucker factorization of a 3rd order tensor Y .

Setting all the matrices of a Tucker decomposition to the identity matrix but the first gives the Tucker-1 decomposition.

Definition 2.9: A rank- R Tucker-1 decomposition of a tensor $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$ produces a matrix $A \in \mathbb{R}^{I_1 \times R}$, and core tensor $B \in \mathbb{R}^{R \times I_2 \times \dots \times I_N}$ such that

$$Y[i_1, \dots, i_N] = \sum_{r=1}^R A[i_1, r]B[r, i_2, \dots, i_N] \quad (6)$$

entry-wise or more compactly,

$$Y = AB = B \times_1 A = \llbracket B; A \rrbracket.$$

Note we extend the usual definition of matrix-matrix multiplication

$$(AB)[i, j] = \sum_{r=1}^R A[i, r]B[r, j]$$

to tensors B in the compact notation for Tucker-1 decomposition $Y = AB$.

More generally, any number of matrices can be set to the identity matrix giving the Tucker- n decomposition.

Definition 2.10: A rank- (R_1, \dots, R_n) Tucker- n decomposition of a tensor $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$ produces n matrices A_1, \dots, A_n , and core tensor $B \in \mathbb{R}^{R_1 \times \dots \times R_n \times I_{n+1} \times \dots \times I_N}$ such that

$$Y[i_1, \dots, i_N] = \sum_{r_1=1}^{R_1} \dots \sum_{r_N=1}^{R_n} A_1[i_1, r_1] \dots A_n[i_N, r_n] B[r_1, \dots, r_n, i_{n+1}, \dots, i_N] \quad (7)$$

entry-wise, or compactly written in the following three ways,

$$\begin{aligned} Y &= B \times_1 A_1 \times_2 \dots \times_n A_n \times_{n+1} \text{id}_{I_{n+1}} \times_{n+2} \dots \times_N \text{id}_{I_N} \\ Y &= B \times_1 A_1 \times_2 \dots \times_n A_n \\ Y &= \llbracket B; A_1, \dots, A_n \rrbracket. \end{aligned}$$

Lastly, if we set the core tensor B to the identity tensor id_R , we obtain the **canonical decomposition/parallel factors** model (CANDECOMP/PARAFAC or CP for short).

Definition 2.11: A rank- R CP decomposition of a tensor $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$ produces N matrices $A_n \in \mathbb{R}^{I_n \times R}$, such that

$$Y[i_1, \dots, i_N] = \sum_{r=1}^R A_1[i_1, r] \cdots A_r[i_N, r] \quad (8)$$

entry-wise. More compactly, this decomposition can be written using the n -mode product, or with double brackets

$$Y = \text{id}_R \times_1 A_1 \times_2 \dots \times_N A_N = \text{id}_R \bigtimes_n A_n = [\![A_1, \dots, A_N]\!].$$

Note CP decomposition is sometimes referred to as Kruskal decomposition, and requires the core only be diagonal (and not necessarily identity) and the factors A_n have normalized columns $\|A_n[:, r]\|_2 = 1$.

Other factorization models are used that combine aspects of CP and Tucker decomposition [13], are specialized for order 3 tensors [14, 15], or provide alternate decomposition models entirely like tensor-trains [16]. But the (full) Tucker, and its special cases Tucker- n , and CP decomposition are most commonly used extensions of the low-rank matrix factorization to tensors. These factorizations are summarized in Table 1.

Table 1: Summary of common tensor factorizations. Here, N is the order of the factorized tensor.

Name	Bracket Notation	n -mode Product	Entry-wise
Tucker	$[\![A_0; A_1, \dots, A_N]\!]$	$A_0 \times_1 A_1 \times_2 \dots \times_N A_N$	Equation 4
Tucker-1	$[\![A_0; A_1]\!]$	$A_0 \times_1 A_1$	Equation 6
Tucker- n	$[\![A_0; A_1, \dots, A_n]\!]$	$A_0 \times_1 A_1 \times_2 \dots \times_n A_n$	Equation 7
CP	$[\![A_1, \dots, A_N]\!]$	$\text{id}_R \times_1 A_1 \times_2 \dots \times_N A_N$	Equation 8

TODO add discussion on other decompositions - high order SVD (see Kolda and Bader) - HOSVD (see Kolda, Shifted power method for computing tensor eigenpairs)

Tensor decompositions are not necessarily unique. It should be clear that scaling one factor by $x \neq 0$ and dividing another by x yields the same original tensor. Furthermore, fibres and slices can be permuted without affecting the the original tensor. Up to these manipulations, for a fixed rank, there exist criteria that ensures their decompositions are unique [13, 17, 18].

2.2.a Representing Tucker Decompositions

There are implemented in `BlockTensorDecomposition.jl` and can be called, for a third order tensor, with `Tucker((B, A1, A2, A3))`, `Tucker1((B, A1))`, and `CPDecomposition((A1, A2, A3))`. These Julia structs store the tensor in its factored form. We could define the contractions for these types and use the common interface provided by `array`, but it turns out we can reconstruct

the whole tensor more efficiently. If the recombined tensor or particular entries are requested, Julia dispatches on the type of decomposition and calls a particular method of array or getindex. The implementations for efficient array construction and index access are provided below.

```
array(T::Tucker) = multifoldl(tucker_contractions(ndims(T)), factors(T))
tucker_contractions(N) = Tuple((G, A) -> nmode_product(G, A, n) for n in 1:N)
```

TODO add getindex method for Tucker type

```
function array(T::Tucker1)
    B, A = factors(T)
    return B ×₁ A
end

function getindex(T::Tucker1, I::Vararg{Int})
    B, A = factors(T)
    i, J... = I # (i, J) = (I[1], I[begin+1:end])
    return (@view A[i, :]) ∙ view(B, :, J...)
end
```

```
array(CPD::CPDecomposition) =
    mapreduce(vector_outer, +, zip((eachcol.(factors(CPD)))...))

vector_outer(v) = reshape(kron(reverse(v)...), length.(v))

getindex(CPD::CPDecomposition, I::Vararg{Int}) =
    sum(reduce(.*, (@view f[i,:]) for (f,i) in zip(factors(CPD), I)))
```

2.3 Tensor rank

- tensor rank
- constrained rank (nonnegative etc.)

The rank of a matrix $Y \in \mathbb{R}^{I \times J}$ can be defined as the smallest $R \in \mathbb{Z}_+$ such that there exists an exact factorization $Y = AB$ for some $A \in \mathbb{R}^{I \times R}$ and $B \in \mathbb{R}^{R \times J}$.

Although this can be extended to higher order tensors, we must specify under which factorization model we are using. For example, the *CP-rank* R of a tensor Y is the smallest such R that omits an exact CP decomposition of Y .

Definition 2.12: The CP rank of a tensor $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$ is the smallest R such that there exist factors $A_n \in \mathbb{R}^{I_n \times R}$ and $Y = [A_1, \dots, A_N]$,

$$\text{rank}_{\text{CP}}(Y) = \min\{R \mid \exists A_n \in \mathbb{R}^{I_n \times R}, n \in [N] \quad \text{s.t.} \quad Y = [A_1, \dots, A_N]\}.$$

In a similar way, we can define the *Tucker-1-rank* R .

Definition 2.13: The Tucker-1 rank of a tensor $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$ is the smallest R such that there exist factors $A \in \mathbb{R}^{I_1 \times R}$ and $B \in \mathbb{R}^{R \times I_2 \times \dots \times I_N}$ where $Y = AB$

$$\text{rank}_{\text{Tucker-1}}(Y) = \min\{R \mid \exists A_n \in \mathbb{R}^{I_n \times R}, B \in \mathbb{R}^{R \times I_2 \times \dots \times I_N} \quad \text{s.t.} \quad Y = AB\}$$

For the Tucker and Tucker- n decompositions, we instead call a particular factorization **a** rank- (R_1, \dots, R_N) Tucker factorization or **a** rank- (R_1, \dots, R_n) Tucker- n factorization, rather than **the** CP- or Tucker-1-rank of a tensor or **the** rank of a matrix.

One reason CP and Tucker-1 only need a single rank R can be explained by considering the case when the order of the tensor $N = 2$ (matrices). The two factorizations become equivalent and are equal to low-rank R matrix factorization $Y = AB$. In fact, Tucker-1 is always equivalent to a low-rank matrix factorization, if you consider a flattening of the tensor to arrange the entries as a matrix.

The idea of tensor rank can be generalized further to constrained rank. These are the smallest rank R such that the factors in the decomposition obey the given set of constraints.

For example, the nonnegative Tucker-1 rank is defined as

$$\text{rank}_{\text{Tucker-1}}^+(Y) = \min\{R \mid \exists A_n \in \mathbb{R}_+^{I_n \times R}, B \in \mathbb{R}_+^{R \times I_2 \times \dots \times I_N} \quad \text{s.t.} \quad Y = AB\}.$$

More restrictive constraints increase the rank of the tensor since there is less freedom in selecting the factors.

Most tensor decomposition algorithms require the rank as input [CITE] since calculating the rank of the tensor can be NP-hard in general [19]. For applications where the rank is not known a priori, a common strategy is to attempt a decomposition for a variety of ranks, and select the model with smallest rank that still achieves good fit between the factorization and the original tensor.

3 Computing Decompositions

- Given a data tensor and a model, how do we fit the model?

Many tensor decompositions algorithms exist in the literature. Usually, they cyclically (or in a random order) update factors until their reconstruction satisfies some convergence criterion. The base algorithm described in Section 3.2 provides flexible framework for wide class of constrained tensor factorization problems. This framework was selected based on empirical observations where it outperforms other similar algorithms, and has also been observed in the literature [10].

3.1 Optimization Problem

- Least squares (can use KL, 1 norm, etc.)

Ideally, we would be given a data tensor Y and decomposition model, and compute an exact factorization of Y into its factors. Because there is often measurement, numerical, or modeling error, an exact factorization of Y for a particular rank may not exist. To over come this, we instead try to fit the model to the data. Let X be the reconstruction of factors A_1, \dots, A_N according to some decomposition for a fixed rank. We assume we know the size of the factors A_1, \dots, A_N and how they are combined to produce a tensor the same size of Y , i.e. the map $g : (A_1, \dots, A_N) \mapsto X$.

There are many loss functions that can be used to determine how close the model X is to the data Y . In principle, any distance or divergence $d(Y, X)$ could be used. We use the L_2 loss or least-squares distance between the tensors $\|X - Y\|_F^2$, but other losses are used for tensor decomposition in practice such as the KL divergence [CITE].

The main optimization we must solve is now given.

Definition 3.1: The constrained least-squares tensor factorization problem is to solve

$$\min_{A_1, \dots, A_N} \frac{1}{2} \|g(A_1, \dots, A_N) - Y\|_F^2 \quad \text{s.t. } (A_1, \dots, A_N) \in \mathcal{C}_1 \times \dots \times \mathcal{C}_N \quad (9)$$

for a given data tensor Y , constraints $\mathcal{C}_1, \dots, \mathcal{C}_N$, and decomposition model g with fixed rank.

Note the problem would have the same solutions as simply using the objective $\|g(A_1, \dots, A_N) - Y\|$ without squaring and dividing by 2. We define the objective in Equation 9 to make computing the function value and gradients faster.

3.2 Base algorithm

- Use Block Coordinate Descent / Alternating Proximal Descent
 - do *not* use alternating least squares (slower for unconstrained problems, no closed form update for general constrained problems)

Let $f(A_1, \dots, A_N) := \frac{1}{2} \|g(A_1, \dots, A_N) - Y\|_F^2$ be the objective function we wish to minimize in Equation 9. Following Xu and Yin [10], the general approach we take to minimize f is to apply block coordinate descent using each factor as a different block. Let A_n^t be the t th iteration of the n th factor, and let

$$f_n^t(A_n) := \frac{1}{2} \|g(A_1^{t+1}, \dots, A_{n-1}^{t+1}, A_n, A_{n+1}^t, \dots, A_N^t) - Y\|_F^2$$

be the (partially updated) objective function at iteration t for factor n .

Given initial factors A_1^0, \dots, A_N^0 , we cycle through the factors $n \in [N]$ and perform the update

$$A_n^{t+1} \leftarrow \arg \min_{A_n \in \mathcal{C}_n} \langle \nabla f_n^t(A_n^t), A_n - A_n^t \rangle + \frac{L_n^t}{2} \|A_n - A_n^t\|_F^2,$$

for $t = 1, 2, \dots$ until some convergence criterion is satisfied (see Section 4.2.a).

This implicit update has the *projected gradient descent* closed form solution for convex constraints \mathcal{C}_n ,

$$A_n^{t+1} \leftarrow P_{\mathcal{C}_n} \left(A_n^t - \frac{1}{L_n^t} \nabla f_n^t(A_n^t) \right). \quad (10)$$

We typically choose L_n^t to be the Lipschitz constant of ∇f_n^t , since it is a sufficient condition to guarantee $f_n^t(A_n^{t+1}) \leq f_n^t(A_n^t)$, but other stepsizes can be used in theory [20 (Sec. 1.2.3)].

?ASIDE? To write ∇f_n^t , we have assumed (block) differentiability of the decomposition model g . In practice, most decompositions are “block-linear” (freeze all factors but one and you have a linear function) and in rare cases are “block-affine”. “block-affine” is enough to ensure f_n^t is convex (i.e. f is “block-convex”) so the updates Equation 10 converge to a Nash equilibrium (block minimizer).

3.2.a High level code

To ensure the code stays flexible, the main algorithm of `BlockTensorDecomposition.jl`, `factorize`, is defined at a very high level.

```

factorize(Y; kwargs...) =
    _factorize(Y; (default_kwargs(Y; kwargs...))...)

"""
Inner level function once keyword arguments are set
"""

function _factorize(Y; kwargs...)
    decomposition, previous, updateprevious!, parameters, updateparameters!,
    update!, stats_data, getstats, converged, kwargs = initialize(Y, kwargs)

    while !converged(stats_data; kwargs...)
        # Usually one cycle of updates through each factor in the decomposition
        update!(decomposition; parameters...)

        # This could be the next stepsize or other info used by update!
        updateparameters!(parameters, decomposition, previous)

        push!(stats_data,
              getstats(decomposition, Y, previous, parameters, stats_data))

        # Update one or two previous iterates. For example, used for momentum
        updateprevious!(previous, parameters, decomposition)
    end

    kwargs = postprocess!(decomposition, Y, previous, parameters, stats_data,
                        updateparameters!, getstats, kwargs)

```

```

    return decomposition, stats_data, kwargs
end

```

The magic of the code is in defining the functions at runtime for a particular decomposition requested, from a reasonable set of default keyword arguments. This is discussed further in Section 4.2.

3.2.b Computing Gradients

- Use Auto diff generally
- But hand-crafted gradients and Lipschitz calculations *can* be faster (e.g. symmetrized slice-wise dot product)

Generally, we can use automatic differentiation on f to compute gradients. Some care needs to be taken otherwise the forward or backwards pass will have to be recompiled every iteration since the factors are updated every iteration.

But for Tucker decompositions, we can compute gradients faster than what an automatic differentiation scheme would give, by taking advantage of symmetry and other computational shortcuts.

Starting with the Tucker-1 decomposition (Definition 2.9), we would like to compute $\nabla_B f(B, A)$ and $\nabla_A f(B, A)$ for $f(B, A) = \frac{1}{2} \|AB - Y\|_F^2$ for a given input Y . We have the gradient

$$\nabla_B f(B, A) = A^\top (AB - Y) = (B \times_1 A - Y) \times_1 A^\top \quad (11)$$

by chain rule, but it is more efficient to calculate the gradient as

$$\nabla_B f(B, A) = (A^\top A)B - A^\top Y = B \times_1 (A^\top A) - Y \times_1 A^\top. \quad (12)$$

²For $A \in \mathbb{R}^{I \times R}$, $B \in \mathbb{R}^{R \times J \times K}$, and $Y \in \mathbb{R}^{I \times J \times K}$, Equation 11 requires

$$\underbrace{2IJKR}_{AB-Y} + \underbrace{IJK(2I-1)}_{A^\top(AB-Y)} \sim 2IJKR + 2I^2JK$$

floating point operations (FLOPS) whereas Equation 12 only uses

$$\underbrace{\frac{R(R+1)}{2}(2I-1)}_{A^\top A} + \underbrace{RJK(2I-1)}_{A^\top Y} + \underbrace{2R^2JK}_{(A^\top A)B-(A^\top Y)} \sim 2IJKR + 2R^2JK + IR^2$$

FLOPS³. So for small ranks $R \ll I$, Equation 12 is cheaper.

A similar story can be said about $\nabla_A f(B, A)$ which is most efficiently computed as

$$\nabla_A f(B, A) = A(B \cdot_1 B) - Y \cdot_1 B.$$

²Seeing Equation 11 and Equation 12 written using the 1-mode product shows how it is “backwards” to normal matrix-matrix multiplication.

³Note we have the smaller factor $R(R+1)/2$ and not the expected R^2 number of entries needed to compute $A^\top A$. The product is a symmetric matrix so only the upper or lower triangle of entries needs to be computed.

! Note

For the family of Tucker decompositions, the objective function f is “block-quadratic” with respect to the factors. This means the gradient with respect to a factor is an affine function of that factor. This is exactly what we see in Equation 12 where B is multiplied by the “slope” $A^\top A$ plus a shift of $-Y \times_1 A^\top$.

The associated implementation with BlockTensorDecomposition.jl is shown below. We define a `make_gradient` which takes the decomposition, factor index n , and data tensor Y , and creates a function that computes the gradient for the same type of decomposition. This lets us manipulate the function that computes the gradient, rather than just the computed gradient.

```
function make_gradient(T::Tucker1, n::Integer, Y::AbstractArray; objective::L2,
kwargs...)
    if n==0 # the core is the zeroth factor
        function gradient0(X::Tucker1; kwargs...)
            (B, A) = factors(X)
            AA = A'A
            YA = Y×_1A'
            grad = B×_1AA - YA
            return grad
        end
        return gradient0
    elseif n==1 # the matrix is the first factor
        function gradient1(X::Tucker1; kwargs...)
            (B, A) = factors(X)
            BB = slicewise_dot(B, B)
            YB = slicewise_dot(Y, B)
            grad = A*BB - YB
            return grad
        end
        return gradient1
    else
        error("No $(n)th factor in Tucker1")
    end
end
```

Similarly, we also have special methods for the Tucker and CP Decomposition.

The gradient with respect to the core for a full Tucker factorization is

$$\nabla_B f(B, A_1, \dots, A_N) = B \bigtimes_n A_n^\top A_n - Y \bigtimes_n A_n^\top,$$

and the gradient with respect to the matrix factor A_n is

$$\nabla_{A_n} f(B, A_1, \dots, A_N) = A_n (\tilde{X}_n \cdot_n \tilde{X}_n) - Y \cdot_n \tilde{X}_n$$

where

$$\tilde{X}_n = \left(B \bigtimes_{m \neq n} A_m \right) = [\![B; A_1, \dots, A_{n-1}, \text{id}_{R_n}, A_{n+1}, \dots, A_N]\!].$$

```

function make_gradient(T::Tucker, n::Integer, Y::AbstractArray; objective::L2,
kwargs...)
    N = ndims(T)
    if n==0 # the core is the zeroth factor
        function gradient_core(X::AbstractTucker; kwargs...)
            B = core(X)
            matrices = matrix_factors(X)
            gram_matrices = map(A -> A'A, matrices) # gram matrices AA = A'A,
            # BB = B'B, ...
            grad = tuckerproduct(B, gram_matrices)
            - tuckerproduct(Y, adjoint.(matrices))
            return grad
        end
        return gradient_core
    elseif n in 1:N # the matrix factors start at m=1
        function gradient_matrix(X::AbstractTucker; kwargs...)
            B = core(X)
            matrices = matrix_factors(X)
            An = factor(X, n)
            Xn = tuckerproduct(B, matrices; exclude=n)
            grad = An * slicewise_dot(Xn, Xn; dims=n)
            - slicewise_dot(Y, Xn; dims=n)
            return grad
        end
        return gradient_matrix
    else
        error("No $(n)th factor in Tucker")
    end
end

```

For the CP Decomposition, we can simply treat the core as $B = \text{id}_R$ and compute the gradient with respect to the matrix factors similarly to the Tucker decomposition:

$$\nabla_{A_n} f(A_1, \dots, A_N) = A_n (\tilde{X}_n \cdot_n \tilde{X}_n) - Y \cdot_n \tilde{X}_n$$

where

$$\tilde{X}_n = \left(\text{id}_R \bigtimes_{m \neq n} A_m \right) = [\![\text{id}_R; A_1, \dots, A_{n-1}, \text{id}_R, A_{n+1}, \dots, A_N]\!].$$

```

function make_gradient(T::CPDecomposition, n::Integer, Y::AbstractArray;
objective::L2, kwargs...)
    N = ndims(T)
    if n in 1:N # the matrix factors start at m=1
        function gradient_matrix(X::AbstractTucker; kwargs...)
            B = core(X)
            matrices = matrix_factors(X)
            A_n = factor(X, n)
            X_n = tuckerproduct(B, matrices; exclude=n)
            grad = A_n * slicewise_dot(X_n, X_n; dims=n)
                - slicewise_dot(Y, X_n; dims=n)
            return grad
        end
        return gradient_matrix
    else
        error("No $(n)th factor in Tucker")
    end
end

```

3.2.c Computing Lipschitz Step-sizes

Similar to automatic differentiation, there exist “automatic Lipschitz” calculations to upper bound the Lipschitz constant of a function [21].

For the family of Tucker decompositions, we can compute the Lipschitz constants of the gradient efficiently similar to how we compute the gradient in Section 3.2.b with the following corollaries of Theorem 2.2.

Corollary 3.1: Let $B \in \mathbb{R}^{R_1 \times \dots \times R_N}$, $A_m \in \mathbb{R}^{I_m \times R_m}$, and $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$. The function

$$f(A) = \frac{1}{2} \| [B; A_1, \dots, A_{n-1}, A, A_{n+1}, \dots, A_N] - Y \|_F^2$$

is quadratic, and L -smooth with constant

$$L_{A_n} = \| \tilde{X}_n \cdot_n \tilde{X}_n \|_{\text{op}}$$

where

$$\tilde{X}_n = \left(B \bigtimes_{m \neq n} A_m \right) = [[B; A_1, \dots, A_{n-1}, \text{id}_{R_n}, A_{n+1}, \dots, A_N]].$$

Proof. The result follows from Theorem 2.1 and Theorem 2.2.

Corollary 3.2: Let $A_n \in \mathbb{R}^{I_n \times R_n}$, and $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$. The function

$$f(B) = \frac{1}{2} \| [B; A_1, \dots, A_N] - Y \|_F^2$$

is quadratic, and L -smooth with constant

$$L_B = \prod_{n=1}^N \|A_n^\top A_n\|_{\text{op}}.$$

Proof. The result follows from Theorem 2.1 and Theorem 2.2.

This yields the following efficient implementations.

! Note

It is tempting to use the identity $\|A^\top A\|_{\text{op}} = \|A\|_{\text{op}}^2$ to calculate the Lipschitz constant without forming $A^\top A$. For tall dense matrices, using this identity is slower and more memory intensive as of Julia 1.11.2. See [LinearAlgebra.jl issue 1185](#) on Github.

```
function make_lipschitz(T::Tucker, n::Integer, Y::AbstractArray; objective::L2,
kwargs...)
    N = ndims(T)
    if n==0 # the core is the zeroth factor
        function lipschitz_core(X::AbstractTucker; kwargs...)
            return prod(A -> opnorm(A'A), matrix_factors(X))
        end
        return lipschitz_core
    elseif n in 1:N # the matrix is the zeroth factor
        function lipschitz_matrix(X::AbstractTucker; kwargs...)
            B = core(X)
            matrices = matrix_factors(X)
            X̃_n = tuckerproduct(B, matrices; exclude=n)
            return opnorm(slicewise_dot(̃X_n, X̃_n; dims=n))
        end
        return lipschitz_matrix
    else
        error("No $(n)th factor in Tucker")
    end
end
```

In the case of Tucker decomposition, the Lipschitz constants simplify to

$$L_B = \|A^\top A\|_{\text{op}}, \quad L_B = \|B \cdot_1 B\|_{\text{op}}.$$

```

function      make_lipschitz(T::Tucker1,      n::Integer,      Y::AbstractArray;
objective::L2, kwargs...)
    if n==0 # the core is the zeroth factor
        function lipschitz0(X::Tucker1; kwargs...)
            A = matrix_factor(X, 1)
            return opnorm(A'A)
        end
        return lipschitz0

    elseif n==1 # the matrix is the zeroth factor
        function lipschitz1(X::Tucker1; kwargs...)
            B = core(X)
            return opnorm(slicewise_dot(B, B))
        end
        return lipschitz1

    else
        error("No $(n)th factor in Tucker1")
    end
end

```

Lastly, for CP decomposition, the Lipschitz constants for the matrices can be calculated similarly to the Tucker decomposition.

```

function      make_lipschitz(T::CPDecomposition,    n::Integer,    Y::AbstractArray;
objective::L2, kwargs...)
    N = ndims(T)
    if n in 1:N
        function lipschitz_matrix(X::CPDecomposition; kwargs...)
            id = core(X)
            matrices = matrix_factors(X)
            X̃_n = tuckerproduct(id, matrices; exclude=n)
            return opnorm(slicewise_dot(X̃_n, X̃_n; dims=n))
        end
        return lipschitz_matrix

    else
        error("No $(n)th factor in CPDecomposition")
    end
end

```

4 Computational Techniques

- As stated, algorithm works
- But can be slow, especially for constrained or large problems

As stated, the algorithm described in Section 3.2 works. It will converge to a solution to our optimization problem and factorize the input tensor. It is worth discussing how the algorithm can be modified to improve convergence to maintain quick convergence for large problems, and what sort of architectural methods are used to allow for maximum flexibility, without over engineering the package.

4.1 For Improving Convergence Speed

There are a few techniques used to assist convergence. Two ideas that are well studied are discussed in this section. They are 1) breaking up the updates into smaller blocks, and 2) using momentum or acceleration. What is perhaps novel is considering the synergy between these two ideas.

Two more techniques are implemented in BlockTensorDecomposition.jl to improve convergence. To the authors knowledge, these are new to tensor factorization, but may or may not be applicable depending on the exact factorization problem or data being studied. For these reasons, these other techniques are discussed separately in Section 5 and Section 6.

4.1.a Sub-block Descent

- Use smaller blocks, but descent in parallel (sub-blocks don't wait for other sub-blocks)
- Can perform this efficiently with a "matrix step-size"

When using block coordinate descent as in Section 3.2, it is natural to treat each factor as its own block. This requires the fewest blocks while ensuring the objective is still convex with respect to each block. We could just as easily use smaller blocks.

In the case of Tucker decomposition, one modification of the update shown in Equation 10 would be to update each column $a_{n,r}$, $r = 1, \dots, R_n$ of the matrix A_n separately. This would be suitable if the constraint that $A_N \in \mathcal{C}_n$ can be broken up further into the constraints $a_{n,r} \in \mathcal{C}_{n,r}$. This is shown in the following update scheme:

$$a_{n,r}^{t+1} \leftarrow P_{\mathcal{C}_{n,r}} \left(a_{n,r}^t - \frac{1}{L_{n,r}^t} \nabla f_{n,r}^t(a_{n,r}^t) \right), \quad (13)$$

where $f_{n,r}^t(a) = \frac{1}{2} \| [B; A_1^{t+1}, \dots, A_{n-1}^{t+1}, A_{n,r}^t(a), A_{n+1}^t, \dots, A_N^t] - Y \|_F^2$ and

$$A_{n,r}^t(a) = \begin{bmatrix} \uparrow & \uparrow & \uparrow & \uparrow & \uparrow \\ a_{n,1}^{t+1} & \cdots & a_{n,r-1}^{t+1} & a & a_{n,r+1}^t & \cdots & a_{n,R_n}^t \\ \downarrow & \downarrow & \downarrow & \downarrow & & & \downarrow \end{bmatrix}. \quad (14)$$

In theory, the block update shown in Equation 13 should be a bit more expensive than using the larger blocks on the matrices A shown in Equation 10, since the gradient needs to be recomputed R_n times for each matrix block n , rather than only computing the gradient once per block n . To get around this, we use the fact that $\nabla f_{n,r}^t(a)$ is the r th column from the gradient $\nabla f_n^t(A)$ where $f_n^t(A) = \frac{1}{2} \| [B; A_1^{t+1}, \dots, A_{n-1}^{t+1}, A, A_{n+1}^t, \dots, A_N^t] - Y \|_F^2$. So we can approximate Equation 13 by first calculating the gradient ∇f_n^t at

$$\hat{A}_n^t = \begin{bmatrix} \uparrow & \uparrow & \uparrow & \uparrow & \uparrow \\ a_{n,1}^t & \cdots & a_{n,r-1}^t & a_{n,r}^t & a_{n,r+1}^t & \cdots & a_{n,R_n}^t \\ \downarrow & & \downarrow & \downarrow & \downarrow & & \downarrow \end{bmatrix}, \quad (15)$$

and then updating each sub-block r according to

$$a_{n,r}^{t+1} \leftarrow P_{\mathcal{C}_{n,r}} \left(a_{n,r}^t - \frac{1}{L_{n,r}^t} \nabla f_n^t(\hat{A}_n^t) \right). \quad (16)$$

Note the difference between Equation 14 and Equation 15 is that we don't use the most recent columns $a_{n,j}$ for $j < r$ in Equation 15.

The update given in Equation 16 can be merged back to an update on the whole block A_n

$$A_n^{t+1} \leftarrow P_{\mathcal{C}_n} \left(A_n^t - \nabla f_n^t(A_n^t) (\hat{L}_n^t)^{-1} \right) \quad (17)$$

where we have the $R_n \times R_n$ diagonal "Lipschitz Matrix"

$$\hat{L}_n^t = \begin{bmatrix} L_{n,1}^t & 0 & 0 \\ 0 & L_{n,2}^t & 0 \\ & \ddots & \vdots \\ 0 & 0 & \cdots & L_{n,R_n}^t \end{bmatrix}.$$

It is not too hard to show that the Lipschitz $L_{n,r}^t$ for $\nabla f_{n,r}^t$ is the Euclidean norm of the r th column⁴ of the matrix $\tilde{X}_n \cdot_n \tilde{X}_n$ from Corollary 3.1,

$$L_{n,r}^t = \|(\tilde{X}_n \cdot_n \tilde{X}_n)[:,r]\|_2.$$

This leads to the following efficient calculation of the Lipschitz matrix in Julia.

```
function diagonal_lipschitz_matrix(T::Tucker, n::Int; kwargs...)
    B = core(T)
    matrices = matrix_factors(T)
    X_n = tuckerproduct(B, matrices; exclude=n)
    return Diagonal_col_norm(slicewise_dot(X_n, X_n; dims=n))
end

Diagonal_col_norm(X) = Diagonal(norm.(eachcol(X)))
```

We can now compare the merged sub-block update Equation 17 to the standard projected gradient descent update shown in Equation 10. The difference is that we calculate a "matrix step-size"

⁴We could have used the r th row of $\tilde{X}_n \cdot_n \tilde{X}_n$ since this matrix is symmetric. Since Julia store matrices in column-major order, many operations that perform column-wise are more efficient than their equivalent row-wise operation.

$\hat{L}_n^t \in \mathbb{R}^{R_n \times R_n}$ rather than a scalar $L_n^t \in \mathbb{R}$. In practice, this leads to an improvement in convergence speed for two reasons.

First, computing the matrix \hat{L}_n^t often faster than the scalar $L_n^t = \|\tilde{X}_n \cdot_n \tilde{X}_n\|_{\text{op}}$. The former only requires calculating the Euclidean norm of R vectors for a total cost of $2R_n^2$ floating point operations (FLOPs), whereas the latter requires the top eigenvalue of $\tilde{X}_n \cdot_n \tilde{X}_n$. This is usually done with a power method or truncated SVD which can be costlier than the flat rate of $2R_n^2$ FLOPs.

Secondly, using the matrix \hat{L}_n^t means columns where $L_{n,r}^t$ is small can take larger descent steps. This is because the largest singular value of $\tilde{X}_n \cdot_n \tilde{X}_n$ is an upper bound on the Euclidean norm of each column: $L_{n,r}^t \leq L_n^t$. Using the scalar Lipschitz L_n^t is equivalent to the diagonal matrix

$$D = \begin{bmatrix} L_n^t & 0 & 0 \\ 0 & L_n^t & 0 \\ & \ddots & \vdots \\ 0 & 0 & \dots & L_n^t \end{bmatrix}$$

in the merged sub-block update shown in Equation 17. So each column of A_n is forced to use the worst case (largest) singular value of $\tilde{X}_n \cdot_n \tilde{X}_n$. In this way, the matrix \hat{L}_n^t acts like a cheap approximate Hessian as if we were doing a quasi-Newton update with step-size 1.

For completeness, we can perform the same merged sub-block update to update the core B . In this case, we obtain the more complicated “Lipschitz tensor” $\hat{L}_B^t \in \mathbb{R}^{(R_1 \times \dots \times R_N)^2}$ defined by

$$\hat{L}_B^t = \hat{L}_{B,1}^t \otimes \dots \otimes \hat{L}_{B,N}^t$$

where each matrix $\hat{L}_{B,n}^t \in \mathbb{R}^{R_n \times R_n}$ is diagonal with non-zero entries

$$L_{B,n}^t[r, r] = \|((A_n^t)^\top A_n^t)[:, r]\|_2.$$

The merged sub-block update for the core becomes

$$B^{t+1} \leftarrow P_{\mathcal{C}_B} \left(B^t - \nabla f_0^t(B^t) \times_B (\hat{L}_B^t)^{-1} \right) \quad (18)$$

with the multiplication

$$\begin{aligned} \nabla f_0^t(B^t) \times_B (\hat{L}_B^t)^{-1} &= \nabla f_0^t(B^t) \bigtimes_n (\hat{L}_{B,n}^t)^{-1} \\ &= [\nabla f_0^t(B^t); (\hat{L}_{B,1}^t)^{-1}, \dots, (\hat{L}_{B,N}^t)^{-1}]. \end{aligned} \quad (19)$$

This should be thought of as normalizing each dimension of the tensor $\nabla f_0^t(B^t)$ so that we can take a unit step-size.

TODO use the multi-mode product in stead of defining a new multiplication \times_B here. I think I'll have the same tensor product issue as before where the indices.

Putting the core and matrices Lipschitz calculations together gives us the following Julia code. Note we store \hat{L}_B^t in factored form as a tuple of diagonal matrices to save space and computation.

TODO Compare to preconditioned decent. See [22–25]

```

function make_block_lipschitz(T::AbstractTucker, n::Integer, Y::AbstractArray;
    objective::L2, kwargs...)
    N = ndims(T)
    if n==0 # the core is the zeroth factor
        function lipschitz_core(X::AbstractTucker; kwargs...)
            return map(A -> Diagonal_col_norm(A'A), matrix_factors(X))
        end # Returns a tuple of diagonal matrices
        return lipschitz_core
    elseif n in 1:N
        function lipschitz_matrix(X::AbstractTucker; kwargs...)
            matrices = matrix_factors(X)
            X̃_n = tuckerproduct(core(X), matrices; exclude=n)
            return Diagonal_col_norm(slicewise_dot(X̃_n, X̃_n; dims=n))
        end
        return lipschitz_matrix
    else
        error("No $(n)th factor in Tucker")
    end
end

```

4.1.b Momentum

- This one is standard
- Use something similar to [10]
- This is compatible with sub-block descent with appropriately defined matrix operations

In practice, we find that extrapolating the iterate based on the prior iterate

$$\hat{A}_n^t \leftarrow A_n^t + \omega_n^t (A_n^t - A_n^{t-1}) \quad (20)$$

for some amount of extrapolation $\omega_n^t \geq 0$ before applying the update Equation 17 greatly improves the speed of descent. This can be thought of as a type of momentum where we continue to move in directions that showed a lot of improvement during the last iteration.

Our selection for ω_n^t follows Xu and Yin’s method for block coordinate descent [10], which is itself inspired by Tseng and Yun’s coordinate gradient descent method [26].

Given a parameter⁵ $\delta \in [0, 1)$, we define the momentum parameters and τ^t and ω_n^t according to the following updates

⁵Usually we pick a number close to 1. For example, we use the default $\delta = 0.9999$.

$$\begin{aligned}
\tau^0 &= 1 \\
\tau^{t+1} &\leftarrow \frac{1}{2} \left(1 + \sqrt{1 + 4(\tau^t)^2} \right) \\
\hat{\omega}^t &\leftarrow \frac{\tau^t - 1}{\tau^{t+1}} \\
\omega_n^t &\leftarrow \min \left(\hat{\omega}^t, \delta \sqrt{\hat{L}_n^{t-1} (\hat{L}_n^t)^{-1}} \right).
\end{aligned} \tag{21}$$

TODO notation is going to get confusing. We use hat/not hat L for the scalar vs matrix/tensor version. But we use hat/not hat ω for the ideal vs clamped momentum.

What is novel with our approach is that we perform this momentum on the Lipschitz matrices and tensors \hat{L}_n^t rather than scalar Lipschitz constant L_n^t . In this way, we should interpret the operations shown in Equation 21 as operating element-wise. This also means the momentum parameter ω_n^t is a matrix or tensor and takes the same shape as \hat{L}_n^t .

In order to perform Equation 20, we use the equivalent but more efficient formulation

$$\hat{A}_n^t \leftarrow A_n^t (\text{id}_{R_n} + \omega_n^t) - A_n^{t-1} \omega_n^t.$$

```

function (U::MomentumUpdate)(X::T; X_last::T, ω, δ, kwargs...) where T
    n = U.n

    L = U.lipschitz(X; kwargs...)
    L_last = U.lipschitz(X_last; kwargs...)
    ω = min.(ω, δ .* √(L_last/L))

    A, A_last = factor(X, n), factor(X_last, n)

    A .= U.combine(A, id + ω)
    A .-= U.combine(A_last, ω)
end

```

In the code above, the momentum stores the factor n it acts on, how to compute the Lipschitz constant, matrix, or tensor, and how to combine (multiply) the constant with the factor. In the case of matrix factors A_n in a Tucker decomposition, this is simply right matrix-matrix multiplication. The core factor B uses \times_B as described in Equation 19.

The parameters of the momentum update are handled separately. This is to treat the momentum update as “apply this update with these parameters”. The parameters τ^t and $\hat{\omega}^t$ are updated by the following function that keeps track of all parameters needed to perform the iteration. In this case, we keep track of what iteration t we are at, the previous iterate, and a few options for the order in which to cycle through and update the blocks.

```

update_τ(τ) = 0.5*(1 + sqrt(1 + 4*τ^2))

function initialize_parameters(decomposition, Y, previous; momentum::Bool,
random_order, recursive_random_order, kwargs...)
    # parameters for the update step are symbol => value pairs
    # held in a dictionary since we may mutate these, e.g. the step-size
    parameters = Dict{Symbol, Any}()

    # General Looping
    parameters[:iteration] = 0
    parameters[:X_last] = previous[begin] # Last iterate
    parameters[:random_order] = random_order
    parameters[:recursive_random_order] = recursive_random_order

    # Momentum
    if momentum
        parameters[:τ_last] = float(1) # need this field to hold Floats, not
Ints
        parameters[:τ] = update_τ(float(1))
        parameters[:ω] = (parameters[:τ_last] - 1) / parameters[:τ]
        parameters[:δ] = kwargs[:δ]
    end

    function updateparameters!(parameters, decomposition, previous)
        parameters[:iteration] += 1
        # parameters[:x_last] = previous[begin]
    # This is commented since parameters[:x_last] already points to previous[begin]

        if momentum
            parameters[:τ_last] = parameters[:τ]
            parameters[:τ] = update_τ(parameters[:τ_last])
            parameters[:ω] = (parameters[:τ_last] - 1) / parameters[:τ]
        end
    end

    return parameters, updateparameters!
end

```

4.1.c Empirical Evidence for Sub-Block Descent and Momentum

To showcase that the combination of these two tricks can speed up convergence, we will benchmark them by factorizing a random 10×10 tensor (a matrix) with rank 3. The Julia code is shown below, and the results are shown in Table 2.

```

using BenchmarkTools
using BlockTensorDecomposition

fact = BlockTensorDecomposition.factorize

```

```

options = (
    :rank => 3,
    :tolerance => (1, 0.03),
    :converged => (GradientNNCone, RelativeError),
    :δ => 0.9,
)
n_subblock_n_momentum(Y) = fact(Y;
    do_subblock_updates=false,
    momentum=false,
    options...
)
y_subblock_n_momentum(Y) = fact(Y;
    do_subblock_updates=true,
    momentum=false,
    options...
)
n_subblock_y_momentum(Y) = fact(Y;
    do_subblock_updates=false,
    momentum=true,
    options...
)
y_subblock_y_momentum(Y) = fact(Y;
    do_subblock_updates=true,
    momentum=true,
    options...
)
I, J = 10, 10
R = 3

@benchmark n_subblock_n_momentum(Y) setup=(Y=Tucker1((I, J), R))
@benchmark n_subblock_y_momentum(Y) setup=(Y=Tucker1((I, J), R))
@benchmark y_subblock_n_momentum(Y) setup=(Y=Tucker1((I, J), R))
@benchmark y_subblock_y_momentum(Y) setup=(Y=Tucker1((I, J), R))

performance_increase(old, new) = (old - new) / new * 100

```

The code `Tucker1((I, J), R)` produces a random $I \times J$ rank- R matrix by generating two matrices $A \in \mathbb{R}^{I \times R}$ and $B \in \mathbb{R}^{R \times J}$ with standard normal entries, and multiplies them together.

Table 2: Summary of median times to factorize a random 10×10 rank-3 matrix under different methods. The performance increase is given by the formula $(\text{old} - \text{new}) / \text{new}$.

	No Momentum	Yes Momentum
No Sub-Block	48.843 ms	45.738 ms (6.7887% faster)
Yes Sub-Block	27.473 ms (77.785% faster)	24.350 ms (100.59% faster)

In Table 2, you can see that having both sub-block descent and momentum yields the fastest factorization. Moreover, the performance increase is *more* than simply the performance increases obtained by exclusively sub-block or momentum alone.⁶ This suggests that there is synergy with these two methods and are best used together.

TODO Repeat this experiment on a less trivial factorization. What I've done above can be done with an SVD in less time. Ideally use a $10 \times 10 \times 10$ Tucker decomposition with rank $2 \times 3 \times 4$.

4.2 For Flexibility

- there are a number of software engineering techniques used
- these help flexibility for hot swapping and a language for making custom...
 - convergence criterion (and having multiple stopping conditions)
 - probing info during the iterations (stats collected at the end)
 - having multiple constraints and ways to enforce them
 - cyclically or partially randomly or fully randomly update factors
- smart enough to apply these in a reasonable order

There are a number of software engineering techniques used to ensure BlockTensorDecomposition.jl is flexible and applicable to a wide range of problems. These enable key algorithmic choices to be hot-swapped and easily compared with each other.

4.2.a Convergence Criteria and Stats

- Can request info about any factor at each outer iteration
- any subset of stats can be the convergence criteria

Some iterative algorithms produce the exact solution of a problem after a finite number of iterations. Generalized minimal residual method (GMRES) is a good example of this [TODO cite!]. Our algorithm, like many others, only converges to the exact solution in the limit as the number of iterations grow. Since we would like a solution in finite time, we must halt the algorithm early.

In finite precision, we can halt the algorithm if we can guarantee the solution is accurate to machine precision. This can often be too strict if convergence is not at a fast enough rate. Furthermore, depending on *why* we are decomposing a tensor, we may want different stats to be within a given a tolerance. BlockTensorDecomposition.jl attempts to solve this issue by defining some standard criteria that can be used to halt the algorithm. These are subtypes of the abstract type `AbstractStat` and are listed below.

⁶The expected performance increase if sub-block descent and momentum where independence would be $(1 + 0.067887)(1 + 0.77785) = 1.89854$ or only 89.854% faster.

```

# X is the decomposition model e.g. Tucker((B, A1, A2, A2)))
# Y is the input tensor we want to decompose
# norm(X) is the Frobenius norm of X

GradientNorm # norm( $\nabla f(X)$ )
GradientNNCone #  $\sqrt{\sum(\text{norm}(\nabla f(A_i)[A_i > 0 \text{ } | \text{ } \nabla f(A_i) < 0])^2 \text{ for } A_i \text{ in factors}(X))}$ 
ObjectiveValue #  $1/2 \text{ norm}(X - Y)^2$ 
ObjectiveRatio #  $\text{norm}(X_{\text{last}} - Y)^2 / \text{norm}(X_{\text{current}} - Y)^2$ 
RelativeError #  $\text{norm}(X - Y) / \text{norm}(Y)$ 
IterateNormDiff #  $\text{norm}(X_{\text{current}} - X_{\text{last}})$ 
IterateRelativeDiff #  $\text{norm}(X_{\text{current}} - X_{\text{last}}) / \text{norm}(X_{\text{last}})$ 

```

Most of these are self explanatory, except perhaps `GradientNNCone`. When performing first order unconstrained optimization, we usually slow down progress when the norm of the gradient is small. In the limit, we expect to converge to a stationary point where the gradient is zero. When performing optimization under nonnegative constraints, and even further restrictions like simplex constraints, it makes more sense to ignore entries of the gradient where X is negative and the gradient is positive since those coordinates of X will not change after they are projected back to the nonnegative constraint.

TODO add theory of negative gradient is in the normal cone of the constraint?

As many or as few of these stats can be used in the call to `factorize`. Their tolerances can also be set independently of each other. The algorithm stops iterating when at least one of the criteria has a value less than its tolerance. As a fail safe, we also define one more type, `Iteration`, which is always active and halts the algorithm when that many iterations have past.

The `AbstractStat` type can also be subtyped to create custom stats that may be used to probe the iterates and diagnose issues.

```

EuclidianStepSize
EuclidianLipschitz
FactorNorms

```

These stats are recorded every iteration in a `DataFrame` and are one of the returns of `factorize`.

Finally, there are two auxiliary stats `PrintStats` and `DisplayDecomposition` which can be used to print all stats or the current decomposition each iteration.

4.2.b Constraints

- one type of update (other than the typical GD update)
- can combine them with composition
 - which is different than projecting onto their intersection!
- Constraint updates combine the constraint with how they are enforced
 - need to go together since there are multiple ways to enforce them e.g. simplex (see next section)

One of the main motivations for developing BlockTensorDecomposition.jl is to solve constrained tensor problems. Other code did not have the expressivity to handle constraints beyond the most common constraints on the factors: nonnegativity and Euclidean normalized columns. To enable flexibility, BlockTensorDecomposition.jl defines

```
abstract type AbstractConstraint <: Function end
```

which has two interfaces. The first treats the constraint like a function

```
(C::AbstractConstraint)(A::AbstractArray)
```

and applies the constraint to an abstract array, and the other interface

```
check(C::AbstractConstraint, A::AbstractArray)
```

checks if the array A satisfies the constraint C. A generic constraint only needs to define these two functions.

```
struct GenericConstraint <: AbstractConstraint
    apply::Function # input a AbstractArray -> mutate it so that `check` would
    return true
    check::Function
end

function (C::GenericConstraint)(D::AbstractArray)
    (C.apply)(D)
end

check(C::GenericConstraint, A::AbstractArray) = (C.check)(A)
```

In general, the function check could be defined as the following,

```
function check(C, A)
    A_copy = copy(A)
    C(A)
    return A ≈ A_copy
end
```

but there is often an easier way to check if a tensor is constrained than to apply the constraint.

Although our basic block projected gradient descent algorithm Equation 10 relies on Euclidean projections to the relevant constraint set, we want to remain flexible and allow for other maps that move an iterate to the constraint set. So each constraint needs to store more than just the constraint itself (the *what*), but also the map from an iterate to the constraint set (the *how*). When prototyping algorithms, it is worth comparing alternate approaches to see if there is a more effi-

cient method to enforce a given constraint (See Section 5 for an example of constraining to a set without Euclidean projections).

The first class of constraints are entry-wise constraints. This is convenient for defining a scalar function that gets applied to all entries of a tensor.

```
struct Entrywise <: AbstractConstraint
    apply::Function
    check::Function
end

function (C::Entrywise)(A::AbstractArray)
    A .= (C.apply).(A)
end

check(C::Entrywise, A::AbstractArray) = all((C.check).(A))
```

Some common examples would be a nonnegative constraint, constraining entries to an interval, or constraining entries to be one or zero.

```
nonnegative! = Entrywise(x -> max(0, x), x ≥ 0)

IntervalConstraint(a, b) = Entrywise(x -> clamp(x, a, b), x -> a ≤ x ≤ b)

binary! = Entrywise(x > 0.5 ? one(x) : zero(x), x -> x in (0, 1))
```

Another class of constraints are normalizations $\mathcal{C}_{\|\cdot\|_a} = \{v \mid \|v\|_a = 1\}$ for some norm $\|\cdot\|_a$. These can be enforced by a (Euclidean) projection

$$\hat{v} \in \arg \min_{u \in \mathcal{C}_{\|\cdot\|_a}} \|u - v\|_2^2,$$

or a scaling (assuming $v \neq 0$)

$$\hat{v} \leftarrow \frac{v}{\|v\|_a}.$$

These operations agree for the Frobenius norm (entry-wise 2-norm), but are different operations in general. In BlockTensorDecomposition.jl, we define these classes as the following.

TODO simplify the following code?

```
abstract type AbstractNormalization <: AbstractConstraint end

struct ProjectedNormalization <: AbstractNormalization
    norm::Function # calculate the norm of an AbstractArray
    projection::Function # mutate an AbstractArray so it has norm == 1
```

```

whats_normalized::Function # what part of the array is normalized
                           # e.g. eachrow, or omit for the entire array
end

ProjectedNormalization(norm, projection; whats_normalized=identityslice) =
    ProjectedNormalization(norm, projection, whats_normalized)

function (P::ProjectedNormalization)(A::AbstractArray)
    whats_normalized_A = P.whats_normalized(A)
    (P.projection).(whats_normalized_A)
end

check(P::ProjectedNormalization, A::AbstractArray) =
    all((P.norm).(P.whats_normalized(A)) .≈ 1)

struct ScaledNormalization{T<:Union{Real,AbstractArray{<:Real},Function}} <:
AbstractNormalization
    norm::Function # calculate the norm of an AbstractArray
    whats_normalized::Function # what part of the array is normalized
    scale::T # the scale that `whats_normalized` should be normalized to
              # Can be a real, array of reals, or function of the input that
              # that returns a real or array of reals
end

ScaledNormalization(norm;whats_normalized=identityslice,scale=1) =
    ScaledNormalization{typeof(scale)}(norm, whats_normalized, scale)

function      (S::ScaledNormalization{T})(A::AbstractArray)           where
{T<:Union{Real,AbstractArray{<:Real}}}
    whats_normalized_A = S.whats_normalized(A)
    A_norm = (S.norm).(whats_normalized_A) ./ S.scale
    whats_normalized_A ./= A_norm
    return A_norm
end

function (S::ScaledNormalization{T})(A::AbstractArray) where {T<:Function}
    whats_normalized_A = S.whats_normalized(A)
    A_norm = (S.norm).(whats_normalized_A) ./ S.scale(A)
    whats_normalized_A ./= A_norm
    return A_norm
end

check(S::ScaledNormalization{<:Union{Real,AbstractArray{<:Real}}}, A::AbstractArray) =
    all((S.norm).(S.whats_normalized(A)) .≈ S.scale)

```

```
check(S::ScaledNormalization{<:Function}, A::AbstractArray) =
    all((S.norm).(S.whats_normalized(A)) .≈ S.scale(A))
```

We can define some common constraints like L_1 normalized through a projection

```
l1normalize! =
    ProjectedNormalization(l1norm, l1project!)
l1normalize_rows! =
    ProjectedNormalization(l1norm, l1project!; whats_normalized=eachrow)
l1normalize_lslices! =
    ProjectedNormalization(l1norm, l1project!;
        whats_normalized=(x -> eachslice(x; dims=1)))
```

or L_∞ normalized by scaling

```
linftyscale_cols! = ScaledNormalization(linftynorm; whats_normalized=eachcol)
```

or even normalize the 3-fibres of a third order tensor on average!

```
l2scale_average12slices! = ScaledNormalization(l2norm;
    whats_normalized=(x -> eachslice(x; dims=1)),
    scale=(A -> size(A, 2)))
```

Constraints can also be composed. This is *not* the same as the intersection of constraints. This is only a handy way to apply multiple constraints in series, but there is no clever logic that interprets the constraints and combines them into a single constraint.

```
struct ComposedConstraint{T<:AbstractConstraint, U<:AbstractConstraint} <:
AbstractConstraint
    outer::T
    inner::U
end

function (C::ComposedConstraint)(A::AbstractArray)
    C.inner(A)
    C.outer(A)
end

check(C::ComposedConstraint, A::AbstractArray) =
    check(C.outer, A) & check(C.inner, A)

Base.:∘(f::AbstractConstraint, g::AbstractConstraint) =
    ComposedConstraint(f, g)
```

This means the following three constraints are all different!

```

l1normalize! ∘ nonnegative!
nonnegative! ∘ l1normalize!
simplex!

```

See Section 5 for a discussion on why it may be advantages to use one of these constraints over the other.

4.2.c BlockUpdate Language

- construct the updates as a list of updates
- very functional programming
- can apply them in sequence or in a random order (or partially random)

To put all these pieces together, we define an abstract type that can be subtyped for the various types of update. This will include gradient descent steps, momentum updates, and constraint enforcing updates.

```

abstract type AbstractUpdate <: Function end

```

A generic update is a function that can take some keyword arguments and mutate an iterate.

```

struct GenericUpdate <: AbstractUpdate
    f::Function
end

(U::GenericUpdate)(x; kwargs...) = U.f(x; kwargs...)

```

The first types of updates are gradient descent updates.

```

abstract type AbstractGradientDescent <: AbstractUpdate end

```

This includes the regular gradient descent update, and also the sub-block descent updates.

```

struct GradientDescent <: AbstractGradientDescent
    n::Integer
    gradient::Function
    step::AbstractStep
end

function (U::GradientDescent)(x; x_last, kwargs...)
    n = U.n
    if checkfrozen(x, n)
        return x
    end
    grad = U.gradient(x; kwargs...)
    # Note we pass a function for grad_last (lazy) so that we only compute it

```

```

if needed for the step
    s = U.step(x; n, x_last, grad, grad_last=(x -> U.gradient(x; kwargs...)),
    kwargs...)
    a = factor(x, n)
    @. a -= s*grad
end

```

The only addition to `BlockGradientDescent` is a function that combines the step with the gradient.

```

struct BlockGradientDescent <: AbstractGradientDescent
    n::Integer
    gradient::Function
    step::AbstractStep
    combine::Function # takes a step (number, matrix, or tensor) and combines
    it with a gradient
end

function (U::BlockGradientDescent)(x; x_last, kwargs...)
    n = U.n
    if checkfrozen(x, n)
        return x
    end
    grad = U.gradient(x; kwargs...)
    # Note we pass a function for grad_last (lazy) so that we only compute it
    if needed for the step
        s = U.step(x; n, x_last, grad, grad_last=(x -> U.gradient(x; kwargs...)),
        kwargs...)
        a = factor(x, n)
        a .-= U.combine(grad, s)
    end
end

```

A separate step type is defined to allow for different types of steps like constant step, Lipschitz, and spectral projected gradient (SPG).

```

abstract type AbstractStep <: Function end

struct LipschitzStep <: AbstractStep
    lipschitz::Function
end

function (step::LipschitzStep)(x; kwargs...)
    L = step.lipschitz(x)
    return L^(-1) # allow for Lipschitz to be a diagonal matrix
end

```

```

function (step::LipschitzStep)(x::Tucker; kwargs...)
    L = step.lipschitz(x)
    if typeof(L) <: Tuple # Currently the only case is when we are updating the
    core of a Tucker factorization
        # Using this condition as a way to tell if it is the
        core we are calculating the constant for
        return map(X -> X^(-1), L)
    else
        return L^(-1) # allow for Lipschitz to be a diagonal matrix
    end
end

```

```

struct ConstantStep <: AbstractStep
    stepsize::Real
end

(step::ConstantStep)(x; kwargs...) = step.stepsize

```

```

struct SPGStep <: AbstractStep
    min::Real
    max::Real
end

SPGStep(;min=le-10, max=le10) = SPGStep(min, max)

# Convert an input of the full decomposition, to a calculation on the nth factor
(step::SPGStep)(x::T; n, x_last::T, grad_last::Function, kwargs...) where {T <:
AbstractDecomposition} =
    step(factor(x,n); x_last=factor(x_last,n), grad_last=grad_last(x_last),
kwargs...)

function (step::SPGStep)(x; grad, x_last, grad_last, stepmin=step.min,
stepmax=step.max, kwargs...)
    s = x - x_last
    y = grad - grad_last
    sy = (s ⋅ y)
    if sy <=0
        return stepmax
    else
        suggested_step = (s ⋅ s) / sy
        return clamp(suggested_step, stepmin, stepmax)
    end
end

```

The gradient and Lipschitz stepsize is calculated by a separate function that gets made on initialization of the factorization algorithm (See Section 3.2.b and Section 3.2.c). This de-couples applying the gradient descent (the *what*), and the computation of the gradient (the *how*) so that

the gradient can be calculated manually (if an efficient method is known), or with automatic differentiation. This also allows other updates to use the same computation code. For example, momentum updates also use the same Lipschitz calculation function. Momentum updates also use the same `combine` function as the sub-block gradient descent updates.

```
struct MomentumUpdate <: AbstractUpdate
    n::Integer
    lipschitz::Function
    combine::Function # How to combine the momentum variable `w` with a factor
    `a`
end

MomentumUpdate(n, lipschitz) = MomentumUpdate(n, lipschitz, (w, a) -> w * a)

function MomentumUpdate(GD::AbstractGradientDescent)
    n, step = GD.n, GD.step
    @assert typeof(step) <: LipschitzStep

    return MomentumUpdate(n, step.lipschitz)
end

function MomentumUpdate(GD::BlockGradientDescent)
    n, step, combine = GD.n, GD.step, GD.combine
    @assert typeof(step) <: LipschitzStep

    return MomentumUpdate(n, step.lipschitz, combine)
end
```

Constraints also get their own abstract subtype of updates.

```
abstract type ConstraintUpdate <: AbstractUpdate end
```

We define a constructor for applying a constraint to a particular factor n . This turns an `AbstractConstraint` into a `ConstraintUpdate`.

```
ConstraintUpdate(n, constraint::GenericConstraint; kwargs...) =
    GenericConstraintUpdate(n, constraint)
ConstraintUpdate(n, constraint::ProjectedNormalization; kwargs...) =
    Projection(n, constraint)
ConstraintUpdate(n, constraint::Entrywise; kwargs...) =
    Projection(n, constraint)

function      ConstraintUpdate(n,           constraint::ScaledNormalization;
skip_rescale=false, whats_rescaled=missing, kwargs...)
    if skip_rescale
        ismissing(whats_rescaled) ||
        isnone(whats_rescaled) ||
```

```

    @warn "skip_rescale=true but whats_rescaled=$whats_rescaled was given.
Overriding to whats_rescaled=nothing"
        return Rescale(n, constraint, nothing)
    else
        return Rescale(n, constraint, whats_rescaled)
    end
end

```

A generic constraint update extracts the factor it needs to constrain, and applies the constraint to that factor.

```

struct GenericConstraintUpdate <: ConstraintUpdate
    n::Integer
    constraint::GenericConstraint
end

check(U::GenericConstraintUpdate,           D::AbstractDecomposition) =
    check(U.constraint, factor(D, U.n))

function (U::GenericConstraintUpdate)(x::T; kwargs...) where T
    n = U.n
    A = factor(x, n)
    U.constraint(A)
    check(U, A) ||
        error("Something went wrong with GenericConstraintUpdate:
$GenericConstraintUpdate")
end

```

Projections follow this pattern closely.

```

struct Projection <: ConstraintUpdate
    n::Integer
    proj::Union{ProjectedNormalization, Entrywise}
end

check(P::Projection, D::AbstractDecomposition) = check(P.proj, factor(D, P.n))

function (U::Projection)(x::T; kwargs...) where T
    n = U.n
    U.proj(factor(x, n))
end

```

A more involved type of constraint update is a rescaled update. This uses a scaled normalization, but moves the weight of the factor to other factors.

For example, if we have three matrix factors A, B, C in a CP decomposition where we want the sum of the entries in A to sum to 1, we can divide A by its sum, and multiple factor B by this

amount. That way the recombined tensor $\llbracket A, B, C \rrbracket$ remains unchanged, but now A satisfies the desired constraint. We could instead multiply both B and C by the square root of the sum to achieve a similar outcome. When it is not specified what is rescaled (`whats_rescaled = missing`), we assume we should multiply every other factor by the geometric mean of the scaling. If we just want to scale the factor but skip any rescaling, we can use `whats_rescaled = nothing`.

```

struct Rescale{T<:Union{Nothing, Missing, Function}} <: ConstraintUpdate
    n::Integer
    scale::ScaledNormalization
    whats_rescaled::T
end

check(S::Rescale, D::AbstractDecomposition) = check(S.scale, factor(D, S.n))

function (U::Rescale{<:Function})(x; kwargs...)
    Fn_scale = U.scale(factor(x, U.n))
    to_scale = U.whats_rescaled(x)
    to_scale .*= Fn_scale
end

(U::Rescale{Nothing})(x; kwargs...) =
    U.scale(factor(x, U.n))

function (U::Rescale{Missing})(x; skip_rescale=false, kwargs...)
    Fn_scale = U.scale(factor(x, U.n))
    x_factors = factors(x)
    N = length(x_factors) - 1

    # Nothing to rescale, so return here
    if N == 0 || skip_rescale
        return nothing
    end

    # Assume we want to evenly rescale all other factors by the Nth root of
    Fn_scale
    scale = geomean(Fn_scale)^(1/N)
    for (i, A) in zip(eachfactorindex(x), x_factors)
        # skip over the factor we just updated
        if i == U.n
            continue
        end
        A .*= scale
    end
end

```

See Section 5 for a more in-depth discussion on when it may be beneficial to use this type of constraint update over a simple projection.

We would like to combine all these updates to execute them in serial. We use a `BlockedUpdate` type to do this. This should be thought of as a list of `AbstractUpdates` that get applied one after another.

```
struct BlockedUpdate <: AbstractUpdate
    updates::Vector{AbstractUpdate}
end
```

! Technical Julia Note

We need the updates to be exactly something of the form `AbstractUpdate[]` since we want to push any type of `AbstractUpdates` such as a `MomentumUpdate` or another `BlockedUpdate`, even if not already present. This means it cannot be `Vector{<:AbstractUpdate}` since a `BlockedUpdate` constructed with only `GradientDescent` would give a `GradientDescent[]` vector and we couldn't push a `MomentumUpdate`. And it cannot be `AbstractVector{AbstractUpdate}` since we may not be able to `insert!` or `push!` into other `AbstractVectors` like `Views`.

We forward many standard methods so that `BlockedUpdates` can behave like usual Julia vectors.

TODO should I actually show all these details?

```
Base.getindex(U::BlockedUpdate, i::Int) = getindex(updates(U), i)
Base.getindex(U::BlockedUpdate, I::Vararg{Int}) = getindex(updates(U), I...)
Base.getindex(U::BlockedUpdate, I) = getindex(updates(U), I) # catch all
Base.firstindex(U::BlockedUpdate) = firstindex(updates(U))
Base.lastindex(U::BlockedUpdate) = lastindex(updates(U))
Base.keys(U::BlockedUpdate) = keys(updates(U))
Base.length(U::BlockedUpdate) = length(updates(U))
Base.iterate(U::BlockedUpdate, state=1) = state > length(U) ? nothing : (U[state], state+1)
Base.filter(f, U::BlockedUpdate) = BlockedUpdate(filter(f, updates(U)))
```

We define how `BlockedUpdates` get applied to a tensor with the following function.

```
function (U::BlockedUpdate)(x::T; recursive_random_order::Bool=false,
random_order::Bool=recursive_random_order, kwargs...) where T
    U_updates = updates(U)
    if random_order
        order = shuffle(eachindex(U_updates))
        U_updates = U_updates[order]
    end

    for update! in U_updates
        update!(x; recursive_random_order, kwargs...)
    end
end
```

```

    # note random_order does not get passed down
end
end

```

The default order the blocks are updated is cyclically through each factor of the decomposition `D::AbstractDecomposition`, in the order of `factors(D)`. For `AbstractTucker` decompositions like Tucker, Tucker-1, and CP, this means starting with the core, followed by the matrix factor for the first dimension, second dimension, and so on.

As an example, this would be the default order of updates for nonnegative CP decomposition on an order 3 tensor.

```

BlockedUpdate(
    MomentumUpdate(1, lipschitz)
    GradientStep(1, gradient, LipschitzStep)
    Projection(1, Entrywise(ReLU, isnonnegative))
    MomentumUpdate(2, lipschitz)
    GradientStep(2, gradient, LipschitzStep)
    Projection(2, Entrywise(ReLU, isnonnegative))
    MomentumUpdate(3, lipschitz)
    GradientStep(3, gradient, LipschitzStep)
    Projection(3, Entrywise(ReLU, isnonnegative))
)

```

The order of updates can be randomized with the `random_order` keyword.

```
X, stats, kwargs = factorize(Y; random_order=true)
```

By default, this will keep momentum steps, gradient steps, and constraint steps for each factor together as a block, in this order.

A possible order of updates could be the following. Note that the updates for each factor are grouped together, but each factor is updated in a random order.

```

BlockedUpdate(
    BlockedUpdate(
        MomentumUpdate(2, lipschitz)
        GradientStep(2, gradient, LipschitzStep)
        Projection(2, Entrywise(ReLU, isnonnegative))
    )
    BlockedUpdate(
        MomentumUpdate(1, lipschitz)
        GradientStep(1, gradient, LipschitzStep)
        Projection(1, Entrywise(ReLU, isnonnegative))
    )
    BlockedUpdate(

```

```

        MomentumUpdate(3, lipschitz)
        GradientStep(3, gradient, LipschitzStep)
        Projection(3, Entrywise(ReLU, isnonnegative))
    )
)

```

For more randomization, use the `recursive_random_order` keyword which will also randomize the order in which the momentum steps, gradient steps, and constraint steps are performed.

```
X, stats, kwargs = factorize(Y; recursive_random_order=true)
```

A possible order of updates could now be the following. The updates for each factor are still grouped together, but the updates within each block appear in a random order.

```

BlockedUpdate(
    BlockedUpdate(
        Projection(2, Entrywise(ReLU, isnonnegative))
        MomentumUpdate(2, lipschitz)
        GradientStep(2, gradient, LipschitzStep)
    )
    BlockedUpdate(
        MomentumUpdate(1, lipschitz)
        Projection(1, Entrywise(ReLU, isnonnegative))
        GradientStep(1, gradient, LipschitzStep)
    )
    BlockedUpdate(
        GradientStep(3, gradient, LipschitzStep)
        Projection(3, Entrywise(ReLU, isnonnegative))
        MomentumUpdate(3, lipschitz)
    )
)

```

The opposite of this would be to keep the outer order of blocks as given, but randomize the order which the updates for each factor gets applied, use the following code.

```
X, stats, kwargs = factorize(Y; recursive_random_order=true, random_order=false,
group_by_factor=true)
```

A possible order of updates could now be the following. Note the order of factors is preserved (1, 2, 3) but the inner `BlockedUpdates` have a random order.

```

BlockedUpdate(
    BlockedUpdate(
        Projection(1, Entrywise(ReLU, isnonnegative))
        MomentumUpdate(1, lipschitz)
    )
)

```

```

        GradientStep(1, gradient, LipschitzStep)
    )
    BlockedUpdate(
        MomentumUpdate(2, lipschitz)
        Projection(2, Entrywise(ReLU, isnonnegative))
        GradientStep(2, gradient, LipschitzStep)
    )
    BlockedUpdate(
        GradientStep(3, gradient, LipschitzStep)
        MomentumUpdate(3, lipschitz)
        Projection(3, Entrywise(ReLU, isnonnegative))
    )
)

```

Note all the previously mentioned options still keeps the various updates for each factor together. For full randomization, use the following code.

```
X,      stats,      kwargs      =      factorize(Y;      recursive_random_order=true,
group_by_factor=false)
```

A possible order of updates could now be the following. Note that every update can appear anywhere in the order.

```

BlockedUpdate(
    Projection(3, Entrywise(ReLU, isnonnegative))
    MomentumUpdate(2, lipschitz)
    GradientStep(2, gradient, LipschitzStep)
    MomentumUpdate(1, lipschitz)
    GradientStep(1, gradient, LipschitzStep)
    Projection(2, Entrywise(ReLU, isnonnegative))
    MomentumUpdate(3, lipschitz)
    MomentumUpdate(2, lipschitz)
    Projection(1, Entrywise(ReLU, isnonnegative))
    GradientStep(3, gradient, LipschitzStep)
)

```

The complete behaviour is summarized in Table 3.

We also use `BlockedUpdate` to handle a composition of constraints.

```

ConstraintUpdate(n, constraint::ComposedConstraint; kwargs...) =
    BlockedUpdate(ConstraintUpdate(n, constraint.inner; kwargs...),
                  ConstraintUpdate(n, constraint.outer; kwargs...))
end

```

The BlockUpdate language becomes especially helpful for automatically inserting additional updates as they are requested. For example, if we want to add momentum to a list of updates, we can call the following function.

```
function add_momentum!(U::BlockedUpdate)
    # Find all the GradientDescent updates
    U_updates = updates(U)
    indexes = findall(u -> typeof(u) <: AbstractGradientDescent, U_updates)

    # insert MomentumUpdates before each GradientDescent
    # do this in reverse order so "i" correctly indexes a GradientDescent
    # as we mutate updates
    for i in reverse(indexes)
        insert!(U_updates, i, MomentumUpdate(U_updates[i]))
    end
end
```

We can also interlace two lists of updates to put updates on the same factor next to each other, or group all updates by the factor they act on.

```
function smart_interlace!(U::BlockedUpdate, other_updates)
    for V in other_updates
        smart_insert!(U::BlockedUpdate, V::AbstractUpdate)
    end
end

smart_interlace!(U::BlockedUpdate, V::BlockedUpdate) = smart_interlace!(U::BlockedUpdate, updates(V))

function smart_insert!(U::BlockedUpdate, V::AbstractUpdate)
    U_updates = updates(U)
    i = findlast(u -> u.n == V.n, U_updates)

    # insert the other update immediately after
    # or if there is no update, push it to the end
    isnone(i) ? push!(U_updates, V) : insert!(U_updates, i+1, V)
end
```

```
function group_by_factor(blockedupdate::BlockedUpdate)
    factor_labels = unique(getproperty(U, :n) for U in blockedupdate)
    updates_by_factor = [filter(U -> U.n == n, blockedupdate) for n in factor_labels]
    return BlockedUpdate(updates_by_factor)
end
```

5 Rescaling to Constrain Tensor Factorization

- for bounded linear constraints
 - first project
 - then rescale to enforce linear constraints
- faster to execute than a projection
- often does not loose progress because of the rescaling (decomposition dependent)

Constraints on factors in a tensor decomposition can arise naturally when modeling physical problems. A common class of constraints is normalizations which restrict a factor or slices of a factor to have unit norm. These are sometimes intersected with interval constraints such as requiring entries to be nonnegative.

With constraints being defined in a flexible manner (see Section 4.2.b), we decided to test conventional wisdom that Euclidean projections are the right kind of map to use when enforcing a constraint. A constraint that came up in recent applications of tensor decomposition to geology [27] was enforcing the 1st mode slices of a tensor (e.g. rows in a matrix) or 3rd mode fibres of a third order tensor to lie in their respective simplex.

For example, the demixing of R probability densities b_1, \dots, b_R from I mixtures y_1, \dots, y_I for $I > R$ can be accomplished with a nonnegative matrix factorization (cite). We have the system of equations

$$\begin{aligned} y_1 &= a_{11} b_1 + a_{12} b_2 + \dots + a_{1R} b_R \\ y_2 &= a_{21} b_1 + a_{22} b_2 + \dots + a_{2R} b_R \\ &\vdots \\ y_I &= a_{I1} b_1 + a_{I2} b_2 + \dots + a_{IR} b_R \end{aligned}$$

with unknown mixing coefficients $a_{i,r}$ and densities b_r . If we can discretize the mixtures y_i , we can rewrite this system as a rank R factorization of the matrix Y where $Y[i, :] = y_i$ and

$$\left[\begin{array}{ccc|c} \leftarrow & y_1^\top & \rightarrow \\ \leftarrow & y_2^\top & \rightarrow \\ \leftarrow & \vdots & \rightarrow \\ \leftarrow & y_I^\top & \rightarrow \end{array} \right] = \left[\begin{array}{cccc} a_{11} & a_{12} & \dots & a_{1R} \\ a_{21} & a_{22} & \dots & a_{2R} \\ \vdots & & & \vdots \\ a_{I1} & a_{I2} & \dots & a_{IR} \end{array} \right] \left[\begin{array}{ccc|c} \leftarrow & b_1^\top & \rightarrow \\ \leftarrow & b_2^\top & \rightarrow \\ \leftarrow & \vdots & \rightarrow \\ \leftarrow & b_R^\top & \rightarrow \end{array} \right]$$

$$Y = AB,$$

for matrices A and B .

For the factorization to remain interpretable, we need to ensure each row b_r of B is a density. This means we would like to constrain each row $B[r, :] = b_r$ to the simplex⁷

⁷This assumes the entries $b_r[j]$ in the discretization of the density b_r represent probabilities or areas under some continuous 1D density function, and not the sample values of the density $b_r(x_j)$. One possible discretization is to take J sample points x_j of a grid on the interval $[x_0, x_J]$ where b_r is supported, and define the entries of the discretization to be $B[r, j] = b_r[j] = b_r(x_j)(x_j - x_{j-1})$. For normalized densities $\int_{x_0}^{x_J} b_r(x) dx = 1$, the sum of the entries $\sum_{j=1}^J b_r[j] \approx 1$ when large enough number of samples J are taken.

$$b_r \in \Delta_J = \left\{ v \in \mathbb{R}_+^J \mid \sum_{j=1}^J v[j] = 1 \right\},$$

which we can write as constraining the matrix B to the simplex

$$B \in \Delta_J^R = \left\{ B \in \mathbb{R}_+^{R \times J} \mid \forall r \in [R], \sum_{j=1}^J B[r, j] = 1 \right\}.$$

Given the rows of B represent densities, to ensure the rows of the reconstructed matrix $\hat{Y} = AB$ are still densities, we need the mixing coefficients to be nonnegative $a_{ir} \geq 0$ and rows to sum to one $\sum_r a_{ir} = 1$. This constrains the matrix A to the simplex

$$A \in \Delta_R^I = \left\{ A \in \mathbb{R}_+^{I \times R} \mid \forall i \in [I], \sum_{r=1}^R A[i, r] = 1 \right\}.$$

The question we investigate in this section is the following: how can we best constrain the factors A and B to their respective simplexes, while performing block gradient decent to minimize the least squared error $\frac{1}{2} \|AB - Y\|_F^2$?

5.1 The two approaches for simplex constraints

To constrain a vector $v \in \mathbb{R}^J$ to the simplex

$$\Delta_J = \left\{ v \in \mathbb{R}_+^J \mid \sum_{j=1}^J v[j] = 1 \right\},$$

we could apply a Euclidean projection

$$v \leftarrow \arg \min_{u \in \Delta_J} \|u - v\|_2^2,$$

or a generalized Kullback-Leibler (KL) divergence projection

$$v \leftarrow \arg \min_{u \in \Delta_J} \sum_j u[j] \log \left(\frac{u[j]}{v[j]} \right) - u[j] + v[j] \quad (22)$$

among other reasonable maps onto Δ_J .

The Euclidean simplex projection can be done with the following implementation of Chen and Ye's algorithm [28]. The essence of the algorithm is to efficiently compute the special $t \in \mathbb{R}$ so that

$$v \leftarrow \max(0, v - t\mathbf{1}) \in \Delta_J. \quad (23)$$

The $\max(0, x)$ function should be understood as operating entrywise on x . In BlockTensorDecomposition, we use the helper $\text{ReLU}(x) = \max(0, x)$ for this function to assist with broadcasting.

```

function projsplx(v)
    J = length(v)

    if J==1 # quick exit for trivial length-1 "vectors" (i.e. scalars)
        return [one(eltype(v))]
    end

    v_sorted = sort(v[:]) # Vectorize/extract input and sort all entries
    j = J - 1
    t = 0 # need to ensure t has scope outside the while loop
    while true
        t = (sum(@view v_sorted[j+1:end]) - 1) / (J-j)
        if t >= v_sorted[j]
            break
        else
            j -= 1
        end

        if j >= 1
            continue
        else # j == 0
            t = (sum(v_sorted) - 1) / J
            break
        end
    end
    return ReLU.(v .- t)
end

```

This is turned into an operation that can mutate v with the following definition,

```

function projsplx!(y)
    y .= projsplx(y)
end

```

and can be turned into a `ProjectedNormalization` (see Section 4.2.b) with the following code.

```

simplex! = ProjectedNormalization(isnonnegative_sumtoone, projsplx!)
isnonnegative_sumtoone(x) = all(isnonnegative, x) && sum(x) ≈ 1

```

The generalized Kullback-Leibler divergence projection as stated in Equation 22 is only well-defined when $v[j] > 0$ for all $j \in [J]$. In this case the solution is given by

$$v \leftarrow \frac{v}{\sum_j v[j]}$$

which is well described by Ducellier et. al. [29 (Sec. 2.1)].

To extend the applicability of this map to any v (when there is at least one positive entry $v[j] > 0$), we can first (Euclidean) project onto the nonnegative orthant \mathbb{R}_+^J ,

$$v \leftarrow \arg \min_{u \in \mathbb{R}_+^J} \|u - v\|_2^2 = \max(0, v),$$

and then apply the divergence projection.⁸ All together, this looks like

$$v \leftarrow \frac{\max(0, v)}{\sum_j \max(0, v[j])}. \quad (24)$$

We will refer to Equation 24 as nonnegative projection and rescaling (NNPR). NNPR has the following implementation in BlockTensorDecomposition.jl.

```
l1scale! ◦ nonnegative!
```

We define the two constraints as the following, using the constraint language from Section 4.2.b.

```
nonnegative! = Entrywise(ReLU, isnonnegative)
l1scale! = ScaledNormalization(l1norm)
l1norm(x) = mapreduce(abs, +, x)
```

5.2 The Rescaling Trick for Matrix Factorization

- Explain that we can move the weight from one matrix to another

Comparing the two methods of constraining a vector to the simplex 1) by Euclidean projection (Equation 23) or 2) nonnegative projection and rescaling (NNPR, Equation 24), the latter offers a few advantages. NNPR is cheaper and conceptually easier to compute. Another advantage of NNPR to tensor factorizations, is its ability to constrain a factor without loosing progress while performing gradient descent.

For example, consider the low rank factorization problem of finding matrices A, B such that $Y = AB$ where you would like the sum of entries in B to be one

$$\min_{A \in \mathbb{R}^{I \times R}, B \in \mathbb{R}^{R \times J}} \frac{1}{2} \|AB - Y\|_F^2 \quad \text{s.t. } B \in \Delta_{R,J} = \left\{ B \in \mathbb{R}_+^{R \times J} \mid \sum_{r,j} B[r, j] = 1 \right\}. \quad (25)$$

The basic alternating projected gradient descent algorithm using a Euclidean projection would be

$$\begin{aligned} A &\leftarrow A - \frac{1}{L_A} \nabla_A f(A, B) \\ B &\leftarrow EP_{\Delta_{R,J}} \left(B - \frac{1}{L_B} \nabla_B f(A, B) \right) \end{aligned}$$

⁸In the unfortunate case where every entry of v is nonpositive, we can fallback to the Euclidean simplex projection.

where $f(A, B) = \frac{1}{2} \|AB - Y\|_F^2$ and $EP_{\Delta_{RJ}}$ is the Euclidean projection onto the simplex Δ_{RJ} . In the event the updated value for B ,

$$B - \frac{1}{L_B} \nabla_B f(A, B) := \hat{B} \in \mathbb{R}_+^{R \times J}$$

is already nonnegative, the objective f at the new point $(A, EP_{\Delta_{RJ}}(\hat{B}))$ could be bigger or smaller than the objective before the Euclidean projection $f(A, \hat{B})$.

If we use the nonnegative projection and rescaling NNPR $_{\Delta_{RJ}}$ instead of the Euclidean projection $EP_{\Delta_{RJ}}$ when \hat{B} is already nonnegative, then

$$\text{NNPR}_{\Delta_{RJ}}(\hat{B}) = \frac{1}{\sum_{rj} B[r, j]} \hat{B} := c^{-1} \hat{B}.$$

This means the objective value f at the point $(c^{-1}A, c\hat{B})$ will be the same as the objective value before the KL divergence projection

$$f(cA, \text{NNPR}_{\Delta_{RJ}}(\hat{B})) = f(cA, c^{-1}\hat{B}) = \frac{1}{2} \|Acc^{-1}B - Y\|_F^2 = \frac{1}{2} \|AB - Y\|_F^2 = f(A, \hat{B}).$$

This suggests the following update may be a useful alternative to the standard projected gradient descent

$$\begin{aligned} A &\leftarrow A - \frac{1}{L_A} \nabla_A f(A, B) \\ B &\leftarrow B - \frac{1}{L_B} \nabla_B f(A, B) \\ c &\leftarrow \sum_{r,j} B[r, j] \\ A &\leftarrow cA \\ B &\leftarrow c^{-1}B. \end{aligned}$$

Of course, it is possible that $\hat{B} = B - \frac{1}{L_B} \nabla_B f(A, B)$ has negative entries. So we use both the nonnegative projection and rescaling part of NNPR in the alternating gradient descent with rescaling update

$$\begin{aligned} A &\leftarrow A - \frac{1}{L_A} \nabla_A f(A, B) \\ B &\leftarrow \max\left(0, B - \frac{1}{L_B} \nabla_B f(A, B)\right) \\ c &\leftarrow \sum_{r,j} B[r, j] \\ A &\leftarrow cA \\ B &\leftarrow c^{-1}B. \end{aligned} \tag{26}$$

This algorithm can be called in `BlockTensorDecomposition.jl` with the following code.

```
options = (
    model=Tucker1,
    constraints=[l1scale! ◦ nonnegative!, noconstraint],
)
decomposition, stats, kwargs = factorize(Y; options...);
```

5.3 Generalizing the Matrix Rescaling Trick

The rescaling trick discussed in Section 5.2 applies more generally to other tensor factorization, other simplex-type constraints, and other `ScaledNormalizations`.

5.3.a Simplex-type constraints

Instead of the matrix factorization problem where $B \in \Delta_{RJ}$ is constrained to the full simplex (Equation 25), we can apply the rescaling trick the problem where the rows of B are constrained to the simplex

$$\min_{\substack{A \in \mathbb{R}^{I \times R} \\ B \in \mathbb{R}^{R \times J}}} \frac{1}{2} \|AB - Y\|_F^2 \quad \text{s.t.} \quad B \in \Delta_J^R = \left\{ B \in \mathbb{R}_+^{R \times J} \mid \forall r \in [R], \sum_{j=1}^J B[r, j] = 1 \right\}. \quad (27)$$

This could make sense in applications where rows of B represent probability densities such as the demixing problem discussed at the start of this section (Section 5).

We can adjust the alternating gradient descent update with NNPR (Equation 26) to the following update

$$\begin{aligned} A &\leftarrow A - \frac{1}{L_A} \nabla_A f(A, B) \\ B &\leftarrow \max\left(0, B - \frac{1}{L_B} \nabla_B f(A, B)\right) \\ C[r, r] &\leftarrow \sum_j B[r, j] \quad (C \in \mathbb{R}^{R \times R} \text{ is diagonal}) \\ A &\leftarrow AC \\ B &\leftarrow C^{-1}B. \end{aligned} \quad (28)$$

It is clear that the objective value would be maintained for any invertible matrix C

$$f(AC, C^{-1}B) = \frac{1}{2} \|ACC^{-1}B - Y\|_F^2 = \frac{1}{2} \|AB - Y\|_F^2 = f(A, B).$$

This algorithm can be called in `BlockTensorDecomposition.jl` with the following code.

```
options = (
    model=Tucker1,
```

```

    constraints=[l1scale_rows! . nonnegative!, noconstraint],
)
decomposition, stats, kwargs = factorize(Y; options...);

```

 Warning

This trick would also work if we wanted the columns of A normalized to the simplex. But it does *not* work when we would like each column of B to be constrained to the simplex. The normalizing matrix C would have to be multiplied to the right of B rather than between A and B . A similar story can be said with the rows of A .

5.3.b Other ScaledNormalization's

This trick can also apply to other normalization constraints. For example, we may want the maximum magnitude of each row of B to one. This could make sense in applications where each row represents a waveform audio file (WAV) which has an audio format that takes values between -1 and 1 . Instead of the Euclidean projection⁹

$$v \leftarrow \max(-1, \min(1, v))$$

onto the infinity ball

$$\mathcal{B}_J(\infty) = \left\{ v \in \mathbb{R}^J \mid \max_{j \in [J]} |v[j]| \leq 1 \right\},$$

we can apply the rescaling¹⁰

$$v \leftarrow \frac{v}{\max_{j \in [J]} |v[j]|}.$$

Applying this principle to the factorization problem

$$\min_{\substack{A \in \mathbb{R}^{I \times R} \\ B \in \mathbb{R}^{R \times J}}} \frac{1}{2} \|AB - Y\|_F^2 \quad \text{s.t.} \quad B \in \mathcal{B}_J^R(\infty) = \left\{ B \in \mathbb{R}^{R \times J} \mid \forall r \in [R], \max_{j \in [J]} |B[r, j]| \leq 1 \right\} \quad (29)$$

gives us the update

⁹In audio processing, this is commonly called “clipping”.

¹⁰In audio processing, this is commonly called “normalizing”. Normalizing is often preferred to clipping since it maintains the perceived audio, but just at a different volume than the original signal. Clipping often introduces undesirable distortion (think of sound from a megaphone).

$$\begin{aligned}
A &\leftarrow A - \frac{1}{L_A} \nabla_A f(A, B) \\
B &\leftarrow B - \frac{1}{L_B} \nabla_B f(A, B) \\
C[r, r] &\leftarrow \max_{j \in [J]} |B[r, j]| \quad (C \in \mathbb{R}^{R \times R} \text{ is diagonal}) \\
A &\leftarrow AC \\
B &\leftarrow C^{-1}B.
\end{aligned} \tag{30}$$

This algorithm can be called in `BlockTensorDecomposition.jl` with the following code.

```

options = (
    model=Tucker1,
    constraints=[linfyscale_rows! ° nonnegative!, noconstraint],
)

decomposition, stats, kwargs = factorize(Y; options...);

```

A similar story can be made about other p -norm balls $\mathcal{B}(p)$ or spheres.

5.3.c Other Tensor Factorizations

The rescaling trick is applicable to other tensor factorizations, but is dependent on the exact model and constraints.

The simplest extension of matrix factorization is the Tucker-1 factorization of an order N tensor $Y = B \times_1 A$.

If we would like the first order slices of B to be constrained to the simplex

$$\begin{aligned}
B[r, :, :] &\in \Delta_{JK} = \left\{ B[r, :, :] \in \mathbb{R}_+^{J \times K} \mid \sum_{j,k} B[r, j, k] = 1 \right\} \\
B &\in \Delta_{JK}^R = \left\{ B \in \mathbb{R}_+^{R \times J \times K} \mid \forall r \in [R], \sum_{j,k} B[r, j, k] = 1 \right\},
\end{aligned}$$

we can solve the problem

$$\min_{\substack{A \in \mathbb{R}^{I \times R} \\ B \in \mathbb{R}^{R \times J \times K}}} \frac{1}{2} \|B \times_1 A - Y\|_F^2 \quad \text{s.t. } B \in \Delta_{JK}^R. \tag{31}$$

by iterating the update

$$\begin{aligned}
A &\leftarrow A - \frac{1}{L_A} \nabla_A f(A, B) \\
B &\leftarrow B - \frac{1}{L_B} \nabla_B f(A, B) \\
C[r, r] &\leftarrow \sum_{j \in [J], k \in [K]} B[r, j, k] \quad (C \in \mathbb{R}^{R \times R} \text{ is diagonal}) \\
A &\leftarrow AC \\
B &\leftarrow B \times_1 C^{-1}.
\end{aligned} \tag{32}$$

This algorithm can be called in `BlockTensorDecomposition.jl` with the following code.

```

options = (
    model=Tucker1,
    constraints=[l1scale_1slices! ∘ nonnegative!, noconstraint],
)

decomposition, stats, kwargs = factorize(Y; options...);

```

A setting where this model applies would be if the first order slices of B represent 2-dimensional probability densities.¹¹

For constraints that constrain an entire factor to some scale, the weight of that factor can be distributed to one or multiple other factors. For example, we may wish to find a CP-decomposition of an order 4-tensor $Y = [[A, B, C, D]]$ where the Frobenius norm of A is 1. In the rescaling step, we could move the norm of A to just the matrix B

$$\begin{aligned}
c &\leftarrow \|A\|_F \\
A &\leftarrow c^{-1}A \\
B &\leftarrow cB
\end{aligned}$$

or all three other factors equally

$$\begin{aligned}
c &\leftarrow \|A\|_F \\
A &\leftarrow c^{-1}A \\
B &\leftarrow c^{1/3}B \\
C &\leftarrow c^{1/3}C \\
D &\leftarrow c^{1/3}D.
\end{aligned}$$

In either case, the recombined tensor remains unchanged

$$[[c^{-1}A, cB, C, D]] = [[c^{-1}A, c^{1/3}B, c^{1/3}C, c^{1/3}D]] = [[A, B, C, D]].$$

¹¹If sampling a 2-dimensional probability density $p_r(x, y)$ on a rectangular grid, entries of B could be interpreted as $B[r, j, k] = p_r(x_j, y_k)(x_j - x_{j-1})(y_k - y_{k-1})$.

The two algorithms can be called in `BlockTensorDecomposition.jl` with the following code.

```

options = (
    model=CPDecomposition,
    constraints=ConstraintUpdate(1, l2scaled!=;
        whats_rescaled=(Y -> factor(Y, 2))), # B is the second factor
)

decomposition, stats, kwargs = factorize(Y; options...);

options = (
    model=CPDecomposition,
    constraints=ConstraintUpdate(1, l2scaled!=),
    # assumes all other factors are rescaled
)

decomposition, stats, kwargs = factorize(Y; options...);

```

5.4 Constraining Multiple Factors

When constraining multiple factors with the rescaling approach, there must be at least one factor that is not constrained with rescaling.

Consider the following matrix factorization $Y = AB$ problem where we want both the columns of A and rows of B to be constrained to the simplex.¹²

$$\begin{aligned}
& \min_{\substack{A \in \mathbb{R}^{I \times R} \\ B \in \mathbb{R}^{R \times J}}} \frac{1}{2} \|AB - Y\|_F^2 \\
& \text{s.t.} \\
& A^\top \in \Delta_I^R = \left\{ A^\top \in \mathbb{R}_+^{R \times I} \mid \forall r \in [R], \sum_i A^\top[r, i] = 1 \right\} \\
& B \in \Delta_J^R = \left\{ B \in \mathbb{R}_+^{R \times J} \mid \forall r \in [R], \sum_j B[r, j] = 1 \right\}
\end{aligned} \tag{33}$$

If we try to rescale both A and B while moving the weights to the other factor, we often observe numerical instability or blow up. To be clear, the following update does not seem to work in practice.

¹²We define the constraint on A in terms of A^\top to be consistent with the simplex constraint defined on B .

$$\begin{aligned}
A &\leftarrow \max\left(0, A - \frac{1}{L_A} \nabla_A f(A, B)\right) \\
C_A[r, r] &\leftarrow \sum_{i \in [I]} A[i, r] \\
A &\leftarrow AC_A^{-1} \\
B &\leftarrow C_A B \\
B &\leftarrow \max\left(0, B - \frac{1}{L_B} \nabla_B f(A, B)\right) \\
C_B[r, r] &\leftarrow \sum_{j \in [J]} B[r, j] \\
A &\leftarrow AC_B \\
B &\leftarrow C_B^{-1} B
\end{aligned}$$

The relevant call in BlockTensorDecomposition would be the following.

```

options = (
    model=Tucker1,
    constraints=[ # B is the 0th factor, A is the 1st factor
        ConstraintUpdate(1, l1scale_cols! ∘ nonnegative!;
            whats_rescaled=(x -> eachrow(factor(x, 0)))),
        ConstraintUpdate(0, l1scale_rows! ∘ nonnegative!;
            whats_rescaled=(x -> eachcol(factor(x, 1)))),
    ],
)

decomposition, stats, kwargs = factorize(Y; options...);

```

Instead, one of the factors can be scaled without moving the weight to the other factor. For example, if we wanted to remove the 4th line $B \leftarrow C_A B$, we can call the following.

```

options = (
    model=Tucker1,
    constraints=[ # B is the 0th factor, A is the 1st factor
        ConstraintUpdate(1, l1scale_cols! ∘ nonnegative!;
            whats_rescaled=nothing),
        ConstraintUpdate(0, l1scale_rows! ∘ nonnegative!;
            whats_rescaled=(x -> eachcol(factor(x, 1)))),
    ],
)

decomposition, stats, kwargs = factorize(Y; options...);

```

This approach seems to work better in practice. The same principle of relaxing how constraints are enforced can be applied to the very similar problem

$$\min_{\substack{A \in \mathbb{R}^{I \times R} \\ B \in \mathbb{R}^{R \times J}}} \frac{1}{2} \|AB - Y\|_F^2$$

s.t.

$$A \in \Delta_R^I = \left\{ A \in \mathbb{R}_+^{I \times R} \mid \forall i \in [I], \sum_r A[i, r] = 1 \right\} \quad (34)$$

$$B \in \Delta_J^R = \left\{ B \in \mathbb{R}_+^{R \times J} \mid \forall r \in [R], \sum_j B[r, j] = 1 \right\},$$

where we want both the rows of A and B to be constrained to the simplex. Here, we cannot move the weights from A to B since there are I rows of A but only R rows of B . Instead, we relax the problem to

$$\min_{\substack{A \in \mathbb{R}^{I \times R} \\ B \in \mathbb{R}^{R \times J}}} \frac{1}{2} \|AB - Y\|_F^2$$

s.t.

$$A \in \mathbb{R}_+^{I \times R} \quad (35)$$

$$B \in \Delta_J^R = \left\{ B \in \mathbb{R}_+^{R \times J} \mid \forall r \in [R], \sum_j B[r, j] = 1 \right\}.$$

The relevant update for this relaxed problem would be

$$\begin{aligned} A &\leftarrow \max\left(0, A - \frac{1}{L_A} \nabla_A f(A, B)\right) \\ B &\leftarrow \max\left(0, B - \frac{1}{L_B} \nabla_B f(A, B)\right) \\ C_B[r, r] &\leftarrow \sum_{j \in [J]} B[r, j] \\ A &\leftarrow AC_B \\ B &\leftarrow C_B^{-1}B \end{aligned}$$

and is called in `BlockTensorDecomposition.jl` with the following code.

```
options = (
    model=Tucker1,
    constraints=[ # B is the 0th factor, A is the 1st factor
        ConstraintUpdate(1, nonnegative!),
        ConstraintUpdate(0, l1scale_rows! ∘ nonnegative!;
            whats_rescaled=(x -> eachcol(factor(x, 1)))),
    ],
)
```

```
decomposition, stats, kwargs = factorize(Y; options...);
```

We justify this relaxation with the following argument. When the rows of Y are in the simplex, we can bound how close the rows of A are to summing to one with Theorem 5.1.

Theorem 5.1 (Closeness of A's rows summing to one): Let $Y \in \Delta_J^I$, $A \in \mathbb{R}_+^{I \times R}$, and $B \in \Delta_J^R$ where

$$\|Y - AB\|_F \leq \epsilon.$$

Then for any row $i \in [I]$, the sum of the row $A[i, :]$ is $\epsilon\sqrt{J}$ close to 1

$$\left| 1 - \sum_{r \in [R]} A[i, r] \right| \leq \epsilon\sqrt{J}.$$

Proof. We have the following inequalities.

$$\begin{aligned} \|Y - AB\|_F &\leq \epsilon \\ \frac{1}{\sqrt{J}} \|Y - AB\|_\infty &\leq \epsilon \quad \left(\frac{1}{\sqrt{J}} \|X\|_\infty \leq \|X\|_2 \leq \|X\|_F \text{ for } X \in \mathbb{R}^{I \times J} \right) \\ \frac{1}{\sqrt{J}} \max_{i \in [I]} \left(\sum_{j \in [J]} |(Y - AB)[i, j]| \right) &\leq \epsilon \\ \max_{i \in [I]} \left(\sum_{j \in [J]} |(Y - AB)[i, j]| \right) &\leq \epsilon\sqrt{J} \end{aligned}$$

So for all rows $i \in [I]$,

$$\begin{aligned} \epsilon\sqrt{J} &\geq \sum_{j \in [J]} |(Y - AB)[i, j]| \\ &\geq \left| \sum_{j \in [J]} (Y - AB)[i, j] \right| \\ &= \left| \sum_{j \in [J]} Y[i, j] - \sum_{j \in [J]} (AB)[i, j] \right| \\ &= \left| 1 - \sum_{j \in [J]} \sum_{r \in [R]} A[i, r]B[r, j] \right| \quad (\text{since } Y \in \Delta_J^I) \\ &= \left| 1 - \sum_{r \in [R]} A[i, r] \left(\sum_{j \in [J]} B[r, j] \right) \right| \\ &= \left| 1 - \sum_{r \in [R]} A[i, r] \right| \quad (\text{since } B \in \Delta_J^R). \end{aligned}$$

Theorem 5.1 implies that solutions to the relaxed problem (Equation 35) are approximate solutions to the problem shown in Equation 34. Moreover, if there exists an exact factorization $Y = AB$ for the relaxed problem, then it is also a solution to the original problem.

5.5 Experiment

To illustrate the advantage of this rescaling trick over Euclidean projection, we will consider solving the problem shown in Equation 34 where $Y \in \Delta_J^I$ and there exists an exact factorization $Y = AB$ with $A \in \Delta_R^I$ and $B \in \Delta_J^R$ using eight different algorithms.

6 Multi-scale

- use a coarse discretization along continuous dimensions
- factorize
- linearly interpolate decomposition to warm start larger decompositions

In many applications [30] [TODO add spatial transcriptomics], the data tensor Y represents a discretization of continuous data. In these cases, we can decide how finely or coarsely to sample the data and effectively have control over the dimension of tensor needing to be factored.

For high quality results, we would like as fine of a discretization as possible. Or if the data is already collected in a discretized form, we would like to incorporate all the data. But smaller tensors are faster to decompose because there are fewer parameters to fit and learn. Most matrix operations like addition, multiplication, and finding their norm are also faster for smaller tensors.

We propose a multiresolution approach inspired by wavelets [31] and multigrid [32] that factorizes the data at progressively finer scales. This can greatly speed up how fast large scale tensors can be factorized.

6.1 Basic Approach

We will start with same example from the beginning of Section 5. Give a tensor $Y \in \mathbb{R}_+^{I \times J}$ representing mixtures of discretized densities, we would like to demix the densities according to the model

$$\begin{bmatrix} \leftarrow & y_1^\top & \rightarrow \\ \leftarrow & y_2^\top & \rightarrow \\ \leftarrow & \vdots & \rightarrow \\ \leftarrow & y_I^\top & \rightarrow \end{bmatrix} = \begin{bmatrix} a_{11} & a_{12} & \dots & a_{1R} \\ a_{21} & a_{22} & \dots & a_{2R} \\ \vdots & & & \vdots \\ a_{I1} & a_{I2} & \dots & a_{IR} \end{bmatrix} \begin{bmatrix} \leftarrow & b_1^\top & \rightarrow \\ \leftarrow & b_2^\top & \rightarrow \\ \leftarrow & \vdots & \rightarrow \\ \leftarrow & b_R^\top & \rightarrow \end{bmatrix}$$

or

$$Y = AB.$$

Notice that the rows of Y and B represent samples of continuous densities, so the size of their second dimension J is arbitrary. Suppose each of the 1-dimensional densities are uniformly discretized on an interval $[a, b]$ with J_s number of points. We use s to represent the scale or spacing of the number of points. For example, $J_1 = J$ would be the finest scale using every point in a

discretization $x_1, x_2, x_3, \dots, x_{J_1}$ with $x_1 = a$ and $x_{J_1} = b$, $J_2 = J_1/2$ would be coarser and use every other point $x_1, x_3, x_5, \dots, x_{J_1-1}$.¹³

The basic approach is to factorize $Y_2 \in \mathbb{R}^{I \times J_2}$ with entries $Y_2[i, j] = y_i(x_{2j-1})$ to obtain $A_2^{T_2} \in \mathbb{R}^{I \times R}$ and $B_{T_2} \in \mathbb{R}^{I \times J_2}$ after T_2 many iterations. We use the factors $A_2^{T_2}$ and $B_{T_2}^{T_2}$ to initialize the factorization of $Y_1 = Y \in \mathbb{R}^{I \times J_1}$. We can initialize $A_1^0 = A_2^{T_2}$ since the size of A is the same at both scales, and repeat every entry of B to initialize B_1^0 with entries $B_1^0[i, j] = B_2^{T_2}[i, \lceil j/2 \rceil]$.

Factorizing Y_2 is faster than factorizing Y_1 since 1) there are fewer parameters to learn¹⁴ and 2) most arithmetic like addition and multiplication, as well as calling other operators like `norm` are faster to compute. This gives us a better initialization for A and B in the factorization of Y_1 than some other random initialization so that fewer iterations are needed at the more expensive finer scale.

6.2 General Approach

The basic approach can be generalized in two ways: the data could be continuous in multiple dimensions, and we can recursively apply this multi-scale approach to progressively refine a very coarse factorization.

Suppose we are given a tensor $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$ where the dimensions I_{n_1}, \dots, I_{n_M} represent a gridded discretization of M -dimensional continuous data.

An example of this setting would be an extension of the example shown in Section 6.1 to higher dimensional distributions. We could consider an order-3 tensor where the horizontal slices correspond to a 2-dimensional discretization of a bivariate density. Entries of the input tensor Y would be given by $Y[i, j, k] = f_i(x_j, y_k)$ for continuous probability density functions $f_i : \mathbb{R}^2 \rightarrow \mathbb{R}_+$ for a 2D grid of points (x_j, y_k) . In this example, the second and third dimensions would be continuous.

If we want to perform a rank- (R_1, \dots, R_N) Tucker decomposition of $Y = [A_0; A_1, \dots, A_N]$, we can initialize the factors with a very coarse factorization $A_{n_m}^s \in \mathbb{R}^{J_{n_m}^s \times R_{n_m}}$ where J^s would be a discretization that uses every 2^s points, or more accurately

$$J_{n_m}^s = 2^{\max(S_m - s, 0)} + 1$$

points in total. We select S_m so that at the finest scale $s = 0$, we have $J_{n_m}^0 = I_{n_m}$. The dimensions that do not represent continuous values will use the full sized factors $A_n^s \in \mathbb{R}^{J_n \times R_n}$ with $J_n = I_n$, and the core will remain $A_0^s \in \mathbb{R}^{R_1 \times \dots \times R_N}$ at all scales s .¹⁵

The aim is to fit the product of the factors $[A_0^s; A_1^s, \dots, A_N^s]$ to a lower resolution version of $Y \in \mathbb{R}^{I_1 \times \dots \times I_N}$, namely $Y^s \in \mathbb{R}^{J_1^s \times \dots \times J_N^s}$, and use the result to initialize a finer version of the factors A . The code for this looks like the following.

¹³We assume J is even here, but we could define $J_2 = (J_1 - 1)/2$ if $J_1 = J$ is odd.

¹⁴Only $(I + J_2)R$ at the coarse scale which is less than $(I + J_1)R$ for the fine scale.

¹⁵This assumes the dimensions of Y where Y is continuous have been discretized as one more than a power of two. The same idea holds with some other discretization plan, but becomes more complicated to express notationally and keep track of the number of points at each scale.

```

function multiscale_factorize(Y; kwargs...)
    scales, kwargs = initialize_scales(Y, kwargs)
    coarsest_scale, finer_scales... = scales

    # Factorize Y at the coarsest scale
    Ys = coarsen(Y, coarsest_scale; kwargs...)
    decomposition, stats, kwargs = factorize(Ys; kwargs...)

    # Factorize Y at progressively finer scales
    for scale in finer_scales
        # Use an interpolated version of the coarse factorization
        # as the initialization
        decomposition = interpolate(decomposition, scale; kwargs...)
        kwargs[:decomposition] = decomposition

        Ys = coarsen(Y, scale; kwargs...)
        decomposition, stats, kwargs = factorize(Ys; kwargs...)
    end
    return decomposition, stats, kwargs
end

```

Straightforward subsampled coarsening and constant interpolating can be used for `coarsen` and `interpolate`, but more sophisticated methods can be used in principle. Since the final solve of `factorize` is on the original sized problem, the choice of coarsening and interpolating only influences the initialization used at this finest scale. Below are examples of the basic coarsening and interpolation methods.

```

function coarsen(Y, scale; dims=1:ndims(A), kwargs...)
    N = ndims(A)

    slice = join((d in dims ? "begin:scale:end" : "begin:end" for d in 1:N),
",")
    Y_coarsened = eval(Meta.parse("Y[$(slice)]"))

    return Y_coarsened
    # Y[(d in dims ? (begin:scale:end) : (begin:end) for d in 1:N)...] does not
    # work since it treats end as length(Y), not the length of just that corresponding
    # dimension
end

```

6.3 Convergence of a Multi-scale Method

- show the multi-scale method has tighter bounds than regular gradient descent for lipschitz data
- this assumes no constraints

7 Conclusion

- all-in-one package
- provide a playground to invent new decompositions
- like auto-diff for factorizations

8 Appendix

8.1 Building the Hessian from two gradients

To build the Hessian from the definition of the gradient, we first extend the gradient to tensor-valued functions. For a function $F : \mathbb{R}^{J_1 \times \dots \times J_N} \rightarrow \mathbb{R}^{I_1 \times \dots \times I_M}$ where

$$F(X) = [f_{i_1 \dots i_M}(X)]$$

is a tensor of scalar functions $f_{i_1 \dots i_M} : T \rightarrow \mathbb{R}$, the gradient of F at X is defined entry-wise as

$$\begin{aligned} \nabla F(X)[i_1, \dots, i_M][j_1, \dots, j_N] &= \nabla f_{i_1 \dots i_M}(X)[j_1, \dots, j_N] \\ &= \frac{\partial f_{i_1 \dots i_M}}{\partial X[j_1, \dots, j_N]}(X). \end{aligned}$$

This treats the gradient at X as a tensor of tensors $\nabla F(X) \in (\mathbb{R}^{I_1 \times \dots \times I_M})^{J_1 \times \dots \times J_N}$. This is naturally isomorphic to a tensor of order $M + N$ with entries

$$\nabla F(X)[i_1, \dots, i_M, j_1, \dots, j_N] = \frac{\partial f_{i_1 \dots i_M}}{\partial X[j_1, \dots, j_N]}(X). \quad (36)$$

So we conclude that the gradient at X of a tensor-valued function F is $\nabla F : \mathbb{R}^{J_1 \times \dots \times J_N} \rightarrow \mathbb{R}^{I_1 \times \dots \times I_M \times J_1 \times \dots \times J_N}$ is given by Equation 36.

We can define the Hessian of a scalar function $f : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}$ at X as $\nabla^2 f(X) = \nabla(\nabla f)(X) : \mathbb{R}^{I_1 \times \dots \times I_N} \rightarrow \mathbb{R}^{(I_1 \times \dots \times I_N)^2}$. The inner nabla ∇ is the gradient of the scalar function f , and the outer nabla ∇ is the gradient of the tensor-valued function ∇f .

This means

$$\nabla^2 f(A)[i_1, \dots, i_N, j_1, \dots, j_N] = \frac{\partial^2 f}{\partial A[j_1, \dots, j_N] \partial A[i_1, \dots, i_N]}(A),$$

but if the function has continuous second derivatives, we can perform the partial derivatives in either order

$$\frac{\partial^2 f}{\partial A[j_1, \dots, j_N] \partial A[i_1, \dots, i_N]}(A) = \frac{\partial^2 f}{\partial A[i_1, \dots, i_N] \partial A[j_1, \dots, j_N]}(A).$$

8.2 Randomizing the order of updates

Table 3: Full description of randomizing the order of updates within a BlockUpdate.

group_by_factor	random_order	recursive_random_order	Description
false	false	false	In the order given
false	false	true	In order given, but randomize how existing blocks are ordered (recursively)
false	true	false	Randomize updates, but keep existing blocks in order
false	true	true	Fully random
true	false	false	In the order given
true	false	true	In order of factors, but updates for each factor a random order
true	true	false	Random order of factors, preserve order of updates within each factor
true	true	true	Almost fully random, but updates for each factor are done together

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