

Leveraging GPU Libraries for Efficient Computation of Bayesian Spatial Assignment Models in R

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Project Background

Developing methods to make use of intrinsic markers (genetic and isotopic signals) for the purpose of inferring migratory connectivity.

- Existing methods are too coarse for most applications
- Large amounts of data are available (>150,000 feather samples from >500 species)
- Genetic assignment methods are based on Wasser, et al. (2004)
- Isotopic assignment methods are based on Wunder, et al. (2005)

Preliminary Data (microsats and $\delta^2\text{H}$):

- Hermit Thrush (*Catharus guttatus*) - 138 Individuals, 14 Locations, 6 Loci, 9-27 Alleles
- Wilson's Warbler (*Wilsonia pusilla*) - 163 Individuals, 8 Locations, 9 Loci, 15-31 Alleles

Allele Frequency Model

For the allele i , from locus l , at location k

$$\mathbf{y}_{l \cdot k} \sim \text{MN}(n_{lk} = \sum_i y_{lik}, \mathbf{f}_{l \cdot k})$$

$$f_{lik} = \frac{\exp(X_{lik})}{\sum_i \exp(X_{lik})} \quad \mathbf{X}_{li} \sim \mathcal{N}(\mathbf{M}_{li}, \mathbf{\Sigma})$$

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Likelihood:

$$\begin{aligned} & \prod_l \prod_k \frac{n_{lk}!}{\prod_i y_{lik}!} \prod_i (f_{lik})^{y_{lik}} \\ & \times \prod_l \prod_i 2\pi^{-r/2} |\Sigma|^{-1/2} \exp \left[-\frac{1}{2} (\mathbf{X}_{li} - \mathbf{M}_{li})' \Sigma^{-1} (\mathbf{X}_{li} - \mathbf{M}_{li}) \right] \\ & \times \pi(\boldsymbol{\theta}) \end{aligned}$$

Implementation

Model fitting and prediction is done via MCMC

- Original implementation in pure C++ with minimal dependencies
- Rewritten using R / C++ using Rcpp(Armadillo)
 - Code closer to matrix notation
 - Transparent use of high performance LAPACK implementations (ATLAS, OpenBLAS, Intel MKL)
- GPU based optimizations were added using CUDA, CUBLAS, and MAGMA libraries
- Cross platform R package scatR (hopefully added to CRAN soon)

Performance

System specs - 4 core Intel i5-2500K, GeForce GTX 460

Software specs - Ubuntu 12.04, ATLAS 3.8.4, Cuda 4.2, Magma 1.1

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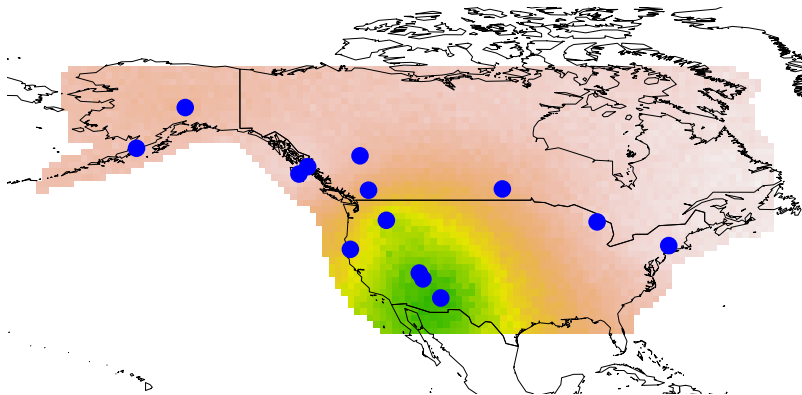
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Performance during prediction is much slower...

100,000 iterations in ~ 2580 seconds (43 mins)
(predictions calculated every 100 iterations)

Not too bad in the greater scheme of things, but we would really like to be able to do cross validation (~ 200 runs per species) ...

Prediction Example



Prediction algorithm details

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To do so we need to draw samples from:

$$\mathbf{X}_p | \mathbf{X}_m \sim \mathcal{N}(\boldsymbol{\mu}_p + \boldsymbol{\Sigma}_{pm} \boldsymbol{\Sigma}_m^{-1} (\mathbf{X}_m - \boldsymbol{\mu}_m), \boldsymbol{\Sigma}_p - \boldsymbol{\Sigma}_{pm} \boldsymbol{\Sigma}_m^{-1} \boldsymbol{\Sigma}_{mp})$$

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Algorithm steps

- 1 Calculate $\boldsymbol{\Sigma}_{pm}$ and $\boldsymbol{\Sigma}_p$
- 2 Calculate $\boldsymbol{\Sigma}_{pm} \boldsymbol{\Sigma}_m^{-1}$
- 3 Calculate $\text{Chol}(\boldsymbol{\Sigma}_p - \boldsymbol{\Sigma}_{pm} \boldsymbol{\Sigma}_m^{-1} \boldsymbol{\Sigma}_{mp})$
- 4 Sample from MVN
- 5 Calculate allele frequencies
- 6 Output results

Prediction algorithm timings

Step	CPU Timing (secs)
1. Covariances	1.02
2. $\Sigma_{21}\Sigma_{11}^{-1}$	0
3. Cholesky	1.15
4. Sample from MVN	0.23
5. Allele Freq	0.14
6. Output	0
Total	2.54

Prediction algorithm timings

Step	CPU Timing (secs)	CPU+GPU (secs)
1. Covariances	1.02	0.05
2. $\Sigma_{21}\Sigma_{11}^{-1}$	0	0
3. Cholesky	1.15	0.23
4. GP Sample	0.23	0.06
5. Allele Freq	0.14	0.14
6. Write	0	0
Total	2.54	0.48

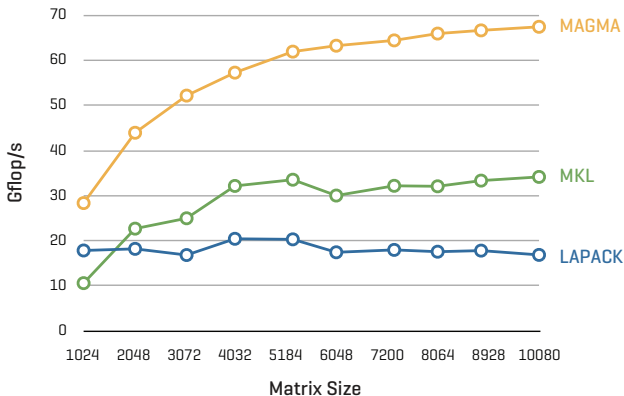
Improving the Cholesky step

Not surprising given Cholesky factorization is $\mathcal{O}(n^3)$ and $n = 3318$.

There isn't a magical solution to this, so we just want to use the fastest possible implementation of the Cholesky decomposition.

- **Intel MKL / OpenBLAS / Eigen** - all (multicore) CPU based with very marginal improvement
- **CUBLAS** - part of NVidia's CUDA toolkit, implements core BLAS functions (but not cholesky)
- **CULA** - proprietary / closed source (dense and sparse) GPU linear algebra library with an expensive license
- **MAGMA** - open source Multicore+GPU dense linear algebra library (CUDA and OpenCL implementations)

MAGMA Performance



Ltaief, H. "A Scalable High Performant Cholesky Factorization for Multicore with GPU Accelerators"

VECTAR'10 Presentation, Berkeley, CA, June 22-25, 2010.

Additional Considerations

- There are costs for moving data on to and off of the GPU
- Once the data is there, may as well do as many calculations as possible
 - Drawing sample from the GP is sped up by performing the matrix multiplication on the GPU
- GPU code is much more verbose / dense

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Armadillo

```
arma::mat tmp = cov12.t() * p.Sinv
```

GPU (CUBLAS)

```
cublasDgemv_v2(  
    p.handle, CUBLAS_OP_T, CUBLAS_OP_N,  
    n_pred, n_known, n_known,  
    &one,  
    p.d_cov12, n_known,  
    p.d_invcov11, n_known,  
    &zero,  
    p.d_tmp, n_pred  
)
```

Improving Covariance calculations

Covariance in our model is assumed to be stationary and isotropic (depend only on distance between locations)

- Elements of the covariance matrix can be calculated independently
- Small scale “embarrassingly parallel” \Rightarrow good candidate for the GPU.
- Implementation is straight forward

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```
__global__ void powered_exponential_kernel(double* dist, double* cov,
                                           const int n, const int nm,
                                           const double sigma2, const double phi,
                                           const double kappa, const double nugget)
{
    int n_threads = blockDim.x * blockDim.x;
    int pos = blockDim.x * blockIdx.x + threadIdx.x;

    for (int i = pos; i < nm; i += n_threads)
        cov[i] = sigma2 * exp(-pow(dist[i] / phi, kappa)) + nugget*(i%n == i/n);
}
```

Summary

Relatively small changes in one function resulted in $\sim 5\times$ improvement

- Cross validation results in days and not weeks
- Started with trying to find an improved Cholesky decomposition, other optimizations followed
- GPU implementation was relatively painless
- Libraries are under active development (read: things can and will break)
- External libraries make package development non-trivial

Questions, Comments?

email : rundel@gmail.com

github : [*http://github.com/rundel/*](http://github.com/rundel/)