LibMints Overview

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LibMints

Provides an interface to compute one- and two-electron integrals with arbitrary angular momentum and substantial flexibility.

With minimal effort rapid implementation is possible, particularly with DF/RI methods.
**LibMints**

LibMints provides a number of C++ classes to the programmer.

For example, fundamental information about the molecular system — atomic coordinates, atomic numbers, symmetry, etc. — is encapsulated in a class called Molecule.
LibMints

A basis set class facilitates the use of multiple basis sets; this is of great utility when dealing with auxiliary basis set methods, such as R12 and DF/RI.

The various types of one- and two-electron integrals that can be computed are all available through a simple, user-friendly interface that is common to all integral types.
Synopsis

• One of the core libraries of Psi4
• Responsible for...
  • managing molecular geometry
  • loading of basis sets
  • computation of integrals
  • providing rudimentary matrix classes
• Interfaces to LIBINT and LIBERD available.
One-Electron Integrals Provided

Well Known One-Electron Integrals
OverlapInt $\langle a|b \rangle$
KineticInt $\langle a| - \frac{1}{2} \nabla^2 | b \rangle$
PotentialInt $\langle a|\frac{1}{|r-C|}|b \rangle$

Property Integrals
AngularMomentumInt
DipoleInt
ElectricFieldInt
ElectrostaticInt
MultipoleInt $\langle a|\mathcal{M}(\mu)|b \rangle$
NablaInt
PseudospectralInt
QuadrupoleInt
Two-Electron Integrals Provided

ERI
ErfComplementERI
ErfERI

Explicitly Correlated Integrals
F12
F12DoubleCommutator
F12G12
F12Squared

CorrelationFactor
FittedSlaterCorrelationFactor
Obtaining the molecule from \texttt{Psi4}

\texttt{Psi4} automatically manages the active molecule for us. To access use the following code.

```cpp
boost::shared_ptr<Molecule> molecule = Process::environment.molecule();
```
Notice

From here on out, I will no longer say boost::
e.g. boost::shared_ptr<>

You will either need to add boost:: yourself, or include the
following near the top of your source file:

```cpp
1 using namespace boost;
```
Loading a basis set

First, a basis set parser object needs to be created.

Second, using your molecule load the necessary basis sets.

```cpp
shared_ptr<BasisSetParser> parser(new Gaussian94BasisSetParser());
shared_ptr<BasisSet> aoBasis = BasisSet::construct(parser, molecule, "BASIS");
```
Information available from a BasisSet

BasisSet contains no symmetry information.

- Number of shells: `nshell()`
- Number of atomic orbitals: `nao()`
- Number of basis functions: `nbf()`

You can also obtain the data for individual shells using

`shell(int shell)`

or

`shell(int center, int shell)`
Obtaining symmetry information for a BasisSet

Symmetry information is obtained through an SOBasisSet object. To construct one, do the following:

```cpp
shared_ptr<SOBasisSet> soBasis(new SOBasisSet(aoBasis, integral));
```

Constructing an integral factory is discussed shortly.
Basis set symmetry information

You obtain symmetry blocking information from an SOBasisSet object:

```cpp
const Dimension dimension = soBasis->dimension();
```

For example, for STO-3G H₂O:

- `aoBasis->nbf()` → 7
- `soBasis->dimension()` → [4, 0, 1, 2]

A `Dimension` object is a helper object for handling matrix dimensionality.
Creating an integral factory

A “factory” is responsible for creating objects.

An “integral factory” is responsible for creating integral objects.

```
shared_ptr<IntegralFactory> integral(
    new IntegralFactory
    (aoBasis, aoBasis, aoBasis, aoBasis)
);
```
Creating one-electron integral objects

Once you have an integral factory creating one-electron integral objects is easy.

```cpp
shared_ptr<OneBodySOInt> sOBI(integral->so_overlap());
shared_ptr<OneBodySOInt> tOBI(integral->so_kinetic());
shared_ptr<OneBodySOInt> vOBI(integral->so_potential());
```
Creating one-electron integral objects

When you’re computing one-electron integrals you’ll need a matrix to store the data in. **LibMints** provides a `MatrixFactory` class to assist you.

Create a matrix factory to help you out and initialize it with the dimensions you obtained from the `SOBasisSet` object:

```cpp
shared_ptr<MatrixFactory> factory(new MatrixFactory);
factory->init_with(dimension, dimension);
```

And then create a matrix:

```cpp
SharedMatrix sMat = factory->create_shared_matrix("Overlap");
```
Compute the overlap integral matrix

Have our previously created integral object compute the integral:

```cpp
// Compute all overlap integrals and store them into sMat
sOBI->compute(sMat);

// Print the matrix
sMat->print();
```
One-electron integrals available

Every entry below is a function available in the IntegralFactory object. Each returns an object that derives from either a OneBodyAOInt or OneBodySOInt class that provides a consistent interface for computing integrals.

<table>
<thead>
<tr>
<th>AO version</th>
<th>SO version</th>
</tr>
</thead>
<tbody>
<tr>
<td>ao_overlap</td>
<td>so_overlap</td>
</tr>
<tr>
<td>ao_kinetic</td>
<td>so_kinetic</td>
</tr>
<tr>
<td>ao_potential</td>
<td>so_potential</td>
</tr>
<tr>
<td>ao_pseudospectral</td>
<td>so_pseudospectral</td>
</tr>
<tr>
<td>ao_dipole</td>
<td>so_dipole</td>
</tr>
<tr>
<td>ao_quadrupole</td>
<td>so_quadrupole</td>
</tr>
<tr>
<td>ao_multipole</td>
<td>so_multipole</td>
</tr>
<tr>
<td>ao_traceless_quadrupole</td>
<td>so_traceless_quadrupole</td>
</tr>
<tr>
<td>ao_nabla</td>
<td>so_nabla</td>
</tr>
<tr>
<td>ao-angular_momentum</td>
<td>so-angular_momentum</td>
</tr>
</tbody>
</table>
Two-electron integrals

\texttt{Psi4} provides the ability to compute AO (no symmetry) and SO (with symmetry) two-electron integrals.

The way to access them is different so let’s cover the AO version first.
Using the integral factory create an ERI object.

// Now, the two−electron integrals
shared_ptr<TwoBodyAOInt> eri(integral−>eri());

We need access to the buffer that the object will put the integrals into:

// buffer will hold the integrals for each shell, as they’re computed
const double *buffer = eri−>buffer();
AO two-electron integrals

**LibMints** provides a couple of handy objects for iterating through only the unique integrals.

```cpp
// The iterator conveniently lets us iterate over functions within shells
AOShellCombinationsIterator shellIter = integral->shells_iterator();

// Let's keep count how many integrals we computed.
unsigned long count=0;

// Use the iterator to loop through the unique shell combinations.
for (shellIter.first(); shellIter.is_done() == false; shellIter.next()) {
```
AO two-electron integrals

Now to compute the integrals.

```c
// Use the iterator to loop through the unique shell combinations.
for (shellIter.first(); shellIter.is_done() == false; shellIter.next()) {
  // Compute quartet
  eri->compute_shell(shellIter);
```
AO two-electron integrals

Work our way through the integrals that we just computed.

```c
// From the quartet get all the integrals
AOIntegralsIterator intIter = shellIter.integrals_iterator();
for (intIter.first(); intIter.is_done() == false; intIter.next()) {
    int p = intIter.i(); int q = intIter.j();
    int r = intIter.k(); int s = intIter.l();
    fprintf(outfile, "\t(%2d %2d | %2d %2d) = %20.15f\n",
            p, q, r, s, buffer[intIter.index()]);
    ++count;
}
fprintf(outfile, "\n\tThere are %d unique integrals\n\n", count);
```
AO two-electron integrals

That little bit of code was enough to compute all the AO two-electron integrals.

What you do with them is up to you.
SO two-electron integrals

The technique used to construct the symmetrized two-electron integrals is different from that of the AO.

We use the C++ technique known as functors.

A functor (or function object) is an C++ class that acts like a function. Functors can be called using the familiar function call syntax, and can yield values and accept parameters just like regular functions.
SO two-electron integrals

Let's first define our functor. Our functor will simply print the integrals out just like our AO integral code before.

```cpp
class ERIPrinter
{
public:
  void operator() (int pabs, int qabs, int rabs, int sabs,
                   int pirrep, int pso,
                   int qirrep, int qso,
                   int rirrep, int rso,
                   int sirrep, int sso,
                   double value)
  {
    fprintf(outfile, "t(%2d %2d | %2d %2d) = %20.10lf\n", 
            pabs, qabs, rabs, sabs, value);
  }
};
```
SO two-electron integrals

Computing SO two-electron integrals takes only a few steps.

First obtain an AO integral object:

```cpp
// 1. Obtain an object that knows how to compute two–electron AO
// integrals.
shared_ptr<TwoBodyAOInt> tb(integral->eri());
```

Create a `TwoBodySOInt` object with your AO integral object.

```cpp
// 2. Create an object that knows how to convert any two–body AO
// integral to SO.
shared_ptr<TwoBodySOInt> eri(new TwoBodySOInt(tb, integral));
```
SO two-electron integrals

Initialize your functor.

```cpp
7  // 3. We to create an instance of our ERIPrinter
8  ERIPrinter printer;
```

Iterator through the shells producing unique SO integrals.

```cpp
9  // 4. Create an SOShellCombinationsIterator to step through the
10 // necessary combinations
11 SOShellCombinationsIterator shellIter(soBasis, soBasis, soBasis, soBasis);
12 for (shellIter.first(); shellIter.is_done() == false; shellIter.next()) {
13    // 5. Call the TwoBodySOInt object to compute integrals giving
14    // it the
15    // instance to our functor.
16    eri->compute_shell(shellIter, printer);
17 }
```
Two-electron integrals available

Every entry below is a function available in the IntegralFactory object. Each returns an object that derives from the TwoBodyAOInt class and provides a consistent interface for computing integrals.

<table>
<thead>
<tr>
<th>AO version</th>
</tr>
</thead>
<tbody>
<tr>
<td>eri</td>
</tr>
<tr>
<td>erd_eri ← Only here for testing purposes.</td>
</tr>
<tr>
<td>erf_eri</td>
</tr>
<tr>
<td>erf_complement_eri</td>
</tr>
<tr>
<td>f12</td>
</tr>
<tr>
<td>f12_squared</td>
</tr>
<tr>
<td>f12g12</td>
</tr>
<tr>
<td>f12_double_commutator</td>
</tr>
</tbody>
</table>
We’ve seen examples of using \texttt{LibMints} to obtain the current molecule from \texttt{Psi4} and computing one- and two- electron integrals.

What else can \texttt{LibMints} do?
What else can LibMINTS do?

- Construct Cartesian displacement SALCs used in finite difference calculations.
- Point group and character tables.
- Integral derivatives.
- Property analysis.
- Also provides a Wavefunction class for theory modules to derive from.
- Various writers for interfacing with other programs. For example, Molden and NBO.
Almost all objects in LibMints are interfaced with Python. This allows for methods for some pretty extraordinary input files. There are a few examples of SCF codes being written in Psi4 input files.
The End.