FastJet 2.4.1 user manual

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Abstract

FastJet provides fast $(N \ln N, N^2)$ implementations of the longitudinally invariant k_t , anti k_t and Cambridge/Aachen jet algorithms for pp collisions, based in part on tools and methods from the Computational Geometry community, as well as a native implementation of the $e^+e^$ k_t algorithm. Further jet algorithms, including most of the other commonly used pp and $e^+e^$ algorithms, can be accessed from the FastJet interface using a plugin mechanism. FastJet also provides ways of determining jet areas.

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1 Introduction

This note documents the FastJet package, which provides efficient geometrically-based implementations [1] for the longitudinally invariant k_t [2, 3], inclusive Cambridge/Aachen [4, 5] and anti- k_t [6] jet algorithms. It also provides access to tools that allow one to determine the areas of individual jets, which is of importance when correcting for underlying event and pileup contamination. Finally external jet algorithms can be accessed through the fastjet interface using a "plugin" facility.

The implementation of the inclusive Cambridge algorithm and of jet areas are new features of version 2 of FastJet; other changes include a new interface,¹ and new algorithmic strategies that can provide a factor of two improvement in speed for events whose number N of particles ~ 10⁴. Choices of recombination schemes and plugins are new features of version 2.1. Version 2.2 introduces a broader set of area measures and the anti- k_t algorithm [6]. A plugin facility allows one to access external jet algorithms through the FastJet interface, and overlays FastJet features such as areas onto the external jet algorithms. Plugins are included for the fortran PxCone code and for the CDF C++ JetClu and MidPoint cone jet algorithms (all infrared unsafe), as well as the recent Seedless Infrared-Safe Cone (SISCone) jet algorithm [7]. Version 2.3 introduced a new build system (GNU autotools), a broader range of areas and tools to help navigate the ClusterSequence Version 2.4 includes the new version 2.0 of SISCone, as well as plugins to the DØ Run II cone, the ATLAS cone, the CMS cone and a range of e^+e^- algorithms, and also further tools to help investigate jet substructure. There is also a wrapper to FastJet allowing one to run SISCone and iterfrom a Fortran program,

The current implementation of the pp sequential-recombination algorithms is restricted to the socalled ΔR distance measure (recommended in [8]), with a choice of recombination schemes. The k_t algorithm has both exclusive [2] and inclusive modes [3], the latter currently being in more widespread use, though the exclusive mode may have physical advantages in certain cases.

2 Quick-start guide

For the impatient, the FastJet package can be set up and run as follows.

• Download the code and the unpack it

```
wget http://www.lpthe.jussieu.fr/~salam/fastjet/repo/fastjet-X.Y.Z.tar.gz
tar zxvf fastjet-X.Y.Z.tar.gz<br>
cd fastjet-X.Y.Z/
```

• Compile and install (choose your own preferred prefix), and when you're done go back to the original directory

```
./configure --prefix='pwd'/../fastjet-install
make
make check
make install
cd ..
```

¹The old one though retained through v2 is deprecated and will be removed in v3

• Now paste the following piece of code into a file called short-example.cc

```
#include "fastjet/ClusterSequence.hh"
 #include <iostream>
 using namespace fastjet;
 using namespace std;
 int main () {
   vector<PseudoJet> particles;
   // an event with two particles:
                                       px py pz
                                                   Е
   particles.push_back( PseudoJet( 100.0, 0, 0, 100.0) );
   particles.push_back( PseudoJet(-100.0, 0, 0, 100.0) );
   // choose a jet definition
   double R = 0.7;
   JetDefinition jet_def(kt_algorithm, R);
   // run the clustering, extract the jets
   ClusterSequence cs(particles, jet_def);
   vector<PseudoJet> jets = cs.inclusive_jets();
   // print the jets
   cout <<
             pt y phi" << endl;</pre>
   for (unsigned i = 0; i < jets.size(); i++) {</pre>
      cout << "jet " << i << ": "<< jets[i].perp() << " "
                     << jets[i].rap() << " " << jets[i].phi() << endl;
   }
 }
• Then compile and run it with
   g++ short-example.cc -o short-example \
       'fastjet-install/bin/fastjet-config --cxxflags' \
       'fastjet-install/bin/fastjet-config --libs'
```

The output will consist of a banner, followed by the lines

pt y phi jet 0: 100 0 3.14159 jet 1: 100 0 0

3 Jet-finding interface

The FastJet code is written in C++. From the point of view of simple usage its interface has a number of similarities to that of KtJet [10], fundamental differences being highlighted below.

All classes are contained in the fastjet namespace. For basic usage, the user is exposed to three main classes:

```
class fastjet::PseudoJet;
class fastjet::JetDefinition;
class fastjet::ClusterSequence;
```

fastjet::PseudoJet provides a jet object with a four-momentum and some internal indices to situate it in the context of a jet-clustering sequence. fastjet::ClusterSequence is the class that carries out jet-clustering and provides access to the final jets.

The class fastjet::JetDefinition contains a specification of how jet clustering is to be performed. Such a class was not present in version 1 of FastJet (nor in KtJet), but was found to be useful to 'contain' the complexity of the interface as new features such as alternative jet algorithms and jet-area determination were introduced.

3.1 fastjet::PseudoJet

All jets, as well as input particles to the clustering (optionally) are fastjet::PseudoJet objects. They can be created using one of the following constructors

```
fastjet::PseudoJet (double px, double py, double pz, double E);
template<class T> fastjet::PseudoJet (const T & some_lorentz_vector);
```

where the second form allows the initialisation to be obtained from any class T that allows subscripting to return the components of the momentum (running from 0...3 in the order p_x, p_y, p_z, E), for example the CLHEP HepLorentzVector class.² It includes the following member functions for accessing the components

```
double E()
                  const ; // returns the energy component
                  const ; // returns the energy component
double e()
                  const ; // returns the x momentum component
double px()
double py()
                  const ; // returns the y momentum component
double pz()
                  const ; // returns the z momentum component
double phi()
                  const ; // returns the azimuthal angle in range 0...2\pi
                  const ; // returns the azimuthal angle in range -\pi \dots \pi
double phi_std()
                  const ; // returns the rapidity
double rap()
double rapidity() const ; // returns the rapidity
double pseudorapidity() const ; // returns the pseudo-rapidity
                  const ; // returns the pseudo-rapidity
double eta()
double kt2()
                  const ; // returns the squared transverse momentum
double perp2()
                  const ; // returns the squared transverse momentum
                  const ; // returns the transverse momentum
double perp()
double m2()
                  const ; // returns squared invariant mass
                  const ; // returns invariant mass ( -\sqrt{-m^2} if m^2 < 0 )
double m()
                  const ; // returns the squared transverse mass = k_t^2 + m^2
double mperp2()
double mperp()
                  const ; // returns the transverse mass
double operator[] (int i) const; // returns component i
double operator() (int i) const; // returns component i
```

/// return a valarray containing the four-momentum (components 0--2

² fastjet::PseudoJet is the analogue of KtJet's KtLorentzVector. A significant difference is that it is not derived from HepLorentzVector (so as to allow compilation even without CLHEP).

```
/// are 3-momentum, component 3 is energy).
valarray<double> four_mom() const;
```

It also allows the user to set and access an index, in case the user wishes to keep track of the identity of a fastjet::PseudoJet object

/// set the user_index, intended to allow the user to label the object
void set_user_index(const int index);

```
/// return the user_index
int user_index() const ;
```

A PseudoJet can be reset with

/// reset the 4-momentum according to the supplied components and /// put the user and history indices back to their default values inline void reset(double px, double py, double pz, double E);

and similarly taking as argument a templated **some_lorentz_vector** or a **PseudoJet** (in the latter case, the user and internal indices are inherited).

Finally, the +, -, * and / operators are defined, with +, - acting on pairs of PseudoJets and *, / acting on a PseudoJet and a double coefficient. Analogous +=, etc., operators, are also defined.

We also provide routines for taking an unsorted vector of fastjet::PseudoJets and returning a sorted vector,

```
/// return a vector of jets sorted into decreasing transverse momentum
vector<fastjet::PseudoJet> sorted_by_pt(const vector<fastjet::PseudoJet> & jets);
/// return a vector of jets sorted into increasing rapidity
vector<fastjet::PseudoJet> sorted_by_rapidity(const vector<fastjet::PseudoJet> & jets);
/// return a vector of jets sorted into decreasing energy
vector<fastjet::PseudoJet> sorted_by_E(const vector<fastjet::PseudoJet> & jets);
```

These will typically be used on the jets returned by fastjet::ClusterSequence.

3.2 fastjet::JetDefinition

The class fastjet::JetDefinition contains a full specification of how to carry out the clustering. According to the Les Houches convention detailed in [11], a 'jet definition' should include the jet algorithm name, its parameters (often the radius R) and the recombination scheme. Its constructor is³

³The v. 2.0 constructor, without the recombination scheme argument, still remains valid.

E_scheme
pt_scheme
pt2_scheme
Et_scheme
Et2_scheme
BIpt_scheme
BIpt2_scheme

Table 1: Members of the **RecombinationScheme** enum; the last two refer to boost-invariant version of the p_t and p_t^2 schemes (as defined in section 5.6).

The jet algorithm is one of the entries of the fastjet::JetAlgorithm enum⁴:

where the ... represent additional values that are present for internal or testing purposes. R specifies the value of R that appears in eqs. (1,2,3,4,5). The recombination scheme should be one of those listed in table 1. If it is omitted the E-scheme is chosen.

For one algorithm, $ee_kt_algorithm$, there is no R parameter, so the constructor is to be called without the R argument:

For the generalised k_t algorithm and its e^+e^- version, one requires R and an extra parameter p, and the following constructor should then be used

fastjet::JetDefinition(fastjet::JetAlgorithm jet_algorithm,

double R, double p, fastjet::RecombinationScheme recomb_scheme = E_scheme, fastjet::Strategy strategy = Best);

If the user calls a constructor with the incorrect number of arguments for the requested jet algorithm, a fastjet::Error() exception will be thrown with an explanatory message.

The default constructor for JetDefinition provides the k_t algorithm with R = 1; the default constructor is there for programming convenience and should not be taken to constitute a default recommendation for physics analyses.

The strategy selects the algorithmic strategy to use while clustering and is an enum of type fastjet::Strategy with potentially interesting values listed in table 2. Nearly all strategies are based on the factorisation of energy and geometrical distance components of the d_{ij} measure [1]. In particular they involve the dynamic maintenance of a nearest-neighbour graph for the geometrical

⁴As of v2.3, the JetAlgorithm name replaces the old JetFinder one, in keeping with the Les Houches convention. Backward compatibility is assured at the user level by a typedef and a doubling of the methods names. Backward compatibility (with versions < 2.3) is however broken for user-written derived classes of ClusterSequence, as the protected variables _default_jet_finder and _jet_finder have been replaced by _default_jet_algorithm and _jet_algorithm.

N2Plain	a plain N^2 algorithm (fastest for $N \lesssim 50$)
N2Tiled	a tiled N^2 algorithm (fastest for $50 \leq N \leq 400$)
N2MinHeapTiled	a tiled N^2 algorithm with a heap for tracking the minimum of
-	d_{ij} (fastest for $400 \leq N \leq 15000$)
NlnN	the Voronoi-based $N \ln N$ algorithm (fastest for $N \gtrsim 15000$)
NlnNCam	based on Chan's $N \ln N$ closest pairs algorithm (fastest for
	$N \gtrsim 6000$), suitable only for the Cambridge jet algorithm
Best	automatic selection of the best of these based on N and R

Table 2: The more interesting of the various algorithmic strategies for clustering. Other strategies are given JetDefinition.hh — note however that strategies not listed in the above table may disappear in future releases. For jet algorithms with spherical distance measures (those whose name starts with "ee_"), only the N2Plain strategy is available.

distances. They apply equally well to any of the internally implemented jet algorithms. The one exception is NlnNCam, which is based on a computational geometry algorithm for dynamic maintenance of closest pairs [12] (rather than the more involved nearest neighbour graph), and is suitable only for the Cambridge algorithm whose distance measure is purely geometrical.

The N2Plain strategy uses a "nearest-neighbour heuristic" [13] approach to maintaining the geometrical nearest-neighbour graph; N2Tiled tiles the $y - \phi$ cylinder to limit the set of points over which nearest-neighbours are searched for,⁵ and N2MinHeapTiled differs only in that it uses an $N \ln N$ (rather than N^2) data structure for maintaining in order the subset of the d_{ij} that involves nearest neighbours. The NlnN strategy uses CGAL's Delaunay Triangulation [15] for the maintenance of the nearest-neighbour graph. Note that $N \ln N$ performance of is an *expected* result, and it holds in practice for the k_t and Cambridge algorithms, while for anti- k_t and generalised- k_t with p < 0, huband-spoke (bicycle-wheel!) type configurations emerge dynamically during the clustering and these break the conditions needed for the expected result to hold (this however has a significant impact only for $N \gtrsim 10^5$).

If strategy is omitted then the Best option is set. Note that the N ranges quoted above for which a given strategy is optimal hold for R = 1; the general R dependence can be significant (and non-trivial), for example for the Cambridge/Aachen jet algorithm with R = 0.4, NlnNCam beats the N2MinHeapTiled strategy only for $N \gtrsim 37000$. While some attempt has been made to account for the R-dependence in the choice of the strategy with the "Best" option, there may exist specific regions of N and R in which a manual choice of strategy can give faster execution. Furthermore the NlnNCam strategy's timings may depend strongly on the size of the cache, and the transitions that have been adopted are based on a cache size of 2 MB. Finally for a given N and R, the optimal strategy may also depend on the event structure.

A textual description of the jet definition can be obtained by a call to the member function

std::string description();

⁵Tiling is a textbook approach in computational geometry, where it is often referred to as bucketing. It has been used also in certain cone jet algorithms, notably at trigger level and in [14].

3.3 fastjet::ClusterSequence

To run the jet clustering, create a fastjet::ClusterSequence object,⁶ through the following constructor

template<class L> fastjet::ClusterSequence

(const std::vector<L> & input_particles, const fastjet::JetDefinition & jet_def);

where input_particles is the vector of initial particles of any type (fastjet::PseudoJet, HepLorentzVector, etc.) that can be used to initialise a fastjet::PseudoJet and jet_def contains the full specification of the clustering (see Section 3.2).

If the user wishes to access inclusive jets, the following member function should be used

```
/// return a vector of all jets (in the sense of the inclusive
/// algorithm) with pt >= ptmin.
vector<fastjet::PseudoJet> inclusive_jets (const double & ptmin = 0.0) const;
```

where **ptmin** may be omitted (then implicitly taking value 0).

There are two ways of accessing exclusive jets,⁷ one where one specifies d_{cut} , the other where one specifies that the clustering is taken to be stopped once it reaches the specified number of jets.

```
/// return a vector of all jets (in the sense of the exclusive
/// algorithm) that would be obtained when running the algorithm
/// with the given dcut.
vector<fastjet::PseudoJet> exclusive_jets (const double & dcut) const;
/// return a vector of all jets when the event is clustered (in the
/// exclusive sense) to exactly njets.
vector<fastjet::PseudoJet> exclusive_jets (const int & njets) const;
```

Note that these two member functions have the same name, and the compiler selects the correct one based on the type of the arguments, as is standard in C++. The fastjet::PseudoJet vectors returned by the above routines can all be sorted with the routines described at the end of section 3.1.

The user may also wish just to obtain information about the number of jets in the exclusive algorithm:

```
/// return the number of jets (in the sense of the exclusive
/// algorithm) that would be obtained when running the algorithm
/// with the given dcut.
int n_exclusive_jets (const double & dcut) const;
```

Another common query is to establish the d_{\min} value associated with merging from n + 1 to n jets. Two member functions are available for determining this:

```
/// return the dmin corresponding to the recombination that went from
/// n+1 to n jets (sometimes known as d_{n n+1}).
double exclusive_dmerge (const int & njets) const;
```

⁶The analogue of KtJet's KtEvent.

⁷In contrast to KtJet the class constructor is the same for the inclusive and exclusive cases. This choice has been made because the clustering sequence is identical in the two cases.

```
/// return the maximum of the dmin encountered during all recombinations
/// up to the one that led to an n-jet final state; identical to
/// exclusive_dmerge, except in cases where the dmin do not increase
/// monotonically.
double exclusive_dmerge_max (const int & njets) const;
```

The first returns the d_{\min} in going from n + 1 to n jets. Occasionally however the d_{\min} value does not increase monotonically during successive mergings and using a d_{cut} smaller than the return value from **exclusive_dmerge** does not guarantee an event with more than **njets** jets. For this reason the second function **exclusive_dmerge_max** is provided — using a d_{cut} below its return value is guaranteed to provide a final state with more than n jets, while using a larger value will return a final state with n or fewer jets.

For e^+e^- collisions, where it is usual to refer to $y_{ij} = d_{ij}/Q^2$ (Q is the total (visible) energy) FastJet provides the following methods:

double exclusive_ymerge (int njets); double exclusive_ymerge_max (int njets); int n_exclusive_jets_ycut (double ycut); std::vector<PseudoJet> exclusive_jets_ycut (double ycut);

which are relevant for use with the $e^+e^- k_t$ algorithm and with the Jade plugin (section 7.4.2).

Finally the user may obtain the constituent particles of a given jet, via

/// return a vector of the particles that make up jet vector<fastjet::PseudoJet> constituents (const fastjet::PseudoJet & jet);

Note that this is a member function of the fastjet::ClusterSequence and not of fastjet::PseudoJet, because jets are only meaningful when referred to a given fastjet::ClusterSequence.⁸ If the user wishes to identify the constituents with the original particles provided to fastjet::ClusterSequence she or he should have set a unique index for each of the original particles with the fastjet::PseudoJet::set_user_index function.

Subjet analysis. To obtain the set of subjets at a specific d_{cut} scale inside a given jet, one may use the following ClusterSequence member function:

If m jets are found, this takes a time $\mathcal{O}(m \ln m)$ (owing to the internal use of a priority queue). Alternatively one may obtain the jet's constituents, cluster them separately and then carry out an **exclusive_jets** analysis on the resulting **ClusterSequence**. The results should be identical. This second method is mandatory if one wishes to identify subjets with an algorithm that differs from the one used to find the original jets.

One can also make use of the following methods, which allow one to follow the merging sequence (and walk back through it):

⁸This is a further respect in which the interface differs from that of KtJet.

Unclustered particles. User-supplied plugin jet algorithms (see section 7) may have the property that not all particles are clustered into jets. In such a case it can be useful to obtain the list of unclustered particles. This can be done as follows:

```
vector<fastjet::PseudoJet> unclustered = clust_seq.unclustered_particles();
```

3.4 Version information

Information on the version of FastJet that is being run can be obtained by making a call to

```
std::string fastjet::fastjet_version_string();
```

(defined in fastjet/JetDefinition.hh). In line with recommendations for other programs in highenergy physics, the user may wish to include this information in publications and plots so as to facilitate reproducibility of the jet-finding.⁹ We recommend also that the main elements of the jet_def.description() be provided, together with citations to the original article that defines the algorithm, as well as to the FastJet paper [1].

4 Example program

For full details see the example program example/fastjet_example.cc that is distributed with the FastJet code. For the reader who is familiar with KtJet [10], the example program is also given as it would be written for KtJet, in the file example/ktjet_example.cc.

A simplified version of the FastJet example program is given below.

```
#include "fastjet/PseudoJet.hh"
#include "fastjet/ClusterSequence.hh"
using namespace std;
```

⁹While we devote significant effort to ensuring that all versions of FastJet give identical, correct results, we are obviously not able to completely guarantee the absence of bugs that might have an effect on the jet finding.

```
int main (int argc, char ** argv) {
  vector<fastjet::PseudoJet> input_particles;
  // read in input particles
  double px, py , pz, E;
  while (cin >> px >> py >> pz >> E) {
    // push fastjet::PseudoJet of (px,py,pz,E) on back of input_particles
    input_particles.push_back(fastjet::PseudoJet(px,py,pz,E));
  }
  // create an object that represents your choice of jet algorithm and
  // its associated parameters
  double Rparam = 1.0;
  fastjet::Strategy strategy = fastjet::Best;
  fastjet::JetDefinition jet_def(fastjet::kt_algorithm, Rparam, strategy);
  // run the jet clustering with the above jet definition
  fastjet::ClusterSequence clust_seq(input_particles, jet_def);
  // extract the inclusive jets with pt > 5 GeV
  double ptmin = 5.0;
  vector<fastjet::PseudoJet> inclusive_jets = clust_seq.inclusive_jets(ptmin);
  // extract the exclusive jets with dcut = 25 GeV<sup>2</sup> and sort them
  // in order of increasing pt
  double dcut = 25.0;
  vector<fastjet::PseudoJet> exclusive_jets = sorted_by_pt(
                                         clust_seq.exclusive_jets(dcut));
  // print out the details for each jet
  for (unsigned int i = 0; i < exclusive_jets.size(); i++) {</pre>
    // get constituents of the jet. NB this is through a member function
    // of clust_seq because that is where the information is held.
    vector<fastjet::PseudoJet> constituents =
                               clust_seq.constituents(exclusive_jets[i])
    printf("%5u %15.8f %15.8f %15.8f %8u\n",
           i, exclusive_jets[i].rap(), exclusive_jets[i].phi(),
           exclusive_jets[i].perp(), constituents.size());
 }
}
```

5 FastJet native jet algorithms

5.1 k_t jet algorithm

The definition of the inclusive k_t jet algorithm that is coded is as follows (and corresponds to [3], modulo small changes of notation):

1. For each pair of particles i, j work out the k_t distance

$$d_{ij} = \min(k_{ti}^2, k_{tj}^2) \,\Delta R_{ij}^2 / R^2 \tag{1}$$

with $\Delta R_{ij}^2 = (y_i - y_j)^2 + (\phi_i - \phi_j)^2$, where k_{ti} , y_i and ϕ_i are the transverse momentum, rapidity and azimuth of particle *i* and *R* is a jet-radius parameter usually taken of order 1; for each parton *i* also work out the beam distance $d_{iB} = k_{ti}^2$.

- 2. Find the minimum d_{\min} of all the d_{ij}, d_{iB} . If d_{\min} is a d_{ij} merge particles *i* and *j* into a single particle, summing their four-momenta (this is *E*-scheme recombination); if it is a d_{iB} then declare particle *i* to be a final jet and remove it from the list.
- 3. Repeat from step 1 until no particles are left.

The exclusive longitudinally invariant k_t jet algorithm [2] is similar except that (a) when a d_{iB} is the smallest value, that particle is considered to become part of the beam jet (i.e. is discarded) and (b) clustering is stopped when all d_{ij} and d_{iB} are above some d_{cut} . In the exclusive mode R is commonly set to 1.

5.2 Cambridge/Aachen jet algorithm

Currently the Cambridge/Aachen jet algorithm is provided only in an inclusive version [5], whose formulation is identical to that of the k_t jet algorithm, except as regards the distance measures, which are:

$$d_{ij} = \Delta R_{ij}^2 / R^2 \,, \tag{2a}$$

$$d_{iB} = 1. (2b)$$

Attempting to extract exclusive jets from the Cambridge/Aachen with a d_{cut} parameter simply provides the set of jets obtained up to the point where all $d_{ij}, d_{iB} > d_{cut}$. Having clustered with some given R, this can actually be an effective way of viewing the event at a smaller radius, $R_{eff} = \sqrt{d_{cut}R}$, thus allowing a single event to be viewed at a continuous range of R_{eff} within a single clustering.

We note that the true exclusive formulation of the Cambridge algorithm [4] instead makes use an auxiliary (k_t) distance measure and 'freezes' pseudojets whose recombination would involve too large a value of the auxiliary distance measure.

5.3 Anti- k_t jet algorithm

This new algorithm, introduced and studied in [6], is defined exactly like the standard k_t algorithm, except for the distance measures which are now given by

$$d_{ij} = \min(1/k_{ti}^2, 1/k_{tj}^2) \,\Delta R_{ij}^2/R^2 \,, \tag{3a}$$

$$d_{iB} = 1/k_{ti}^2 \,. \tag{3b}$$

While being a sequential recombination algorithm like k_t and Cambridge/Aachen, the anti- k_t algorithm behaves in some sense like a 'perfect' cone algorithm, in that its hard jets are exactly circular on the y- ϕ cylinder [6].

5.4 Generalised k_t jet algorithm

The "generalised k_t " algorithm again follows the same procedure, but depends on an additional continuous parameter p, with has the following distance measure:

$$d_{ij} = \min(k_{ti}^{2p}, k_{tj}^{2p}) \,\Delta R_{ij}^2 / R^2 \,, \tag{4a}$$

$$d_{iB} = k_{ti}^{2p} \,. \tag{4b}$$

For specific values of p, it reduces to one or other of the algorithms list above, k_t (p = 1), Cambridge/Aachen (p = 0) and anti- k_t (p = -1).

5.5 Generalised k_t algorithm for e^+e^- collisions

FastJet also provides native implementations of clustering algorithms in spherical coordinates (specifically for e^+e^- collisions) along the lines of the original k_t algorithms [9], but extended in analogy with the generalised pp algorithm of [6] and section 5.4. We define the two following distances:

$$d_{ij} = \min(E_i^{2p}, E_j^{2p}) \frac{(1 - \cos \theta_{ij})}{(1 - \cos R)},$$
(5a)

$$d_{iB} = E_i^{2p} \,, \tag{5b}$$

for a general value of p and R. At a given stage of the clustering sequence, if a d_{ij} is smallest then i and j are recombined, while if a d_{iB} is smallest then i is called an "inclusive jet".

For values of $R \leq \pi$ in eq. (5), the generalised $e^+e^- k_t$ algorithm behaves in analogy with the pp algorithms: when an object is at an angle $\theta_{iX} > R$ from all other objects X then it forms an inclusive jet. With the choice p = -1 this provides a simple, infrared and collinear safe way of obtaining a cone-like algorithm for e^+e^- collisions, since hard well-separated jets have a circular profile on the 3D sphere, with opening half-angle R.

If one imagines a (complex) value of R such that $(1 - \cos R) > 2$, then the d_{iB} will be smallest only if the event consists of a single particle, and thus with the additional choice of p = 1 the clustering sequence will correspond to that of the $e^+e^- k_t$ algorithm [9], often referred to also as the Durham algorithm, which has a single distance:

$$d_{ij} = 2\min(E_i^{2p}, E_j^{2p})(1 - \cos\theta_{ij}).$$
(6)

Note the difference in normalisation between the d_{ij} in eqs. (5) and (6), and the fact in neither case have we normalised to the total energy Q in the event, contrary to the convention adopted originally in [9] (where the distance measure was called y_{ij}).

5.6 Recombination schemes

When merging particles in step 2 of the clustering procedure, one must specify how to combine the momenta. The simplest procedure (E-scheme) simply adds the four-vectors. This has been advocated as a standard in [8], and is the default option in FastJet.

Other schemes for pp collisions. Other schemes provided by earlier k_t -clustering implementations [10] are the p_t , p_t^2 , E_t and E_t^2 schemes. They all incorporate a 'preprocessing' stage to make the initial momenta massless (rescaling the energy to be equal to the 3-momentum for the p_t and p_t^2 schemes, rescaling to the 3-momentum to be equal to the energy in the E_t and E_t^2 schemes). Then for all schemes the recombination p_r of p_i and p_j is a massless 4-vector satisfying

$$p_{t,r} = p_{t,i} + p_{t,j}$$
, (7a)

$$\phi_r = (w_i \phi_i + w_j \phi_j) / (w_i + w_j), \qquad (7b)$$

$$y_r = (w_i y_i + w_j y_j) / (w_i + w_j),$$
 (7c)

where w_i is $p_{t,i}$ for the p_t and E_t schemes, and is $p_{t,i}^2$ for the p_t^2 and E_t^2 schemes.

Note that for massive particles the schemes defined in the previous paragraph are not invariant under longitudinal boosts. We therefore also introduce boost-invariant p_t and p_t^2 schemes, which are identical to the normal p_t and p_t^2 schemes, except that they omit the preprocessing stage.

Other schemes for e^+e^- collisions. On request, we may in the future provide dedicated schemes for e^+e^- collisions.

User-defined schemes. The user may define their own, custom recombination schemes, as described in Appendix A.

6 Jet areas

Since a jet is made up of only a finite number of particles, one needs a specific definition in order to make its area (i.e. the surface in the y- ϕ plane over which it extends) an unambiguous concept. Three definitions of area have been proposed in [16] and they are implemented in FastJet:

- Active areas add a uniform background of extremely soft massless 'ghost' particles to the event and allow them to participate in the clustering. The area of a given jet is proportional to the number of ghosts it contains. Because the ghosts are extremely soft (and sensible jet algorithms are infrared safe), the presence of the ghosts does not affect the set of user particles that end up in a given jet.
- Passive areas are defined as follows. One adds a single randomly placed ghost at a time to the event. One examines which jet (if any) the ghost ends up in. One repeats the procedure many times and the passive area of a jet is then proportional to the probability of it containing the ghost.
- The Voronoi area of a jet is the sum of the Voronoi areas of its constituent particles. The Voronoi area of a particle is obtained by determining the Voronoi diagram for the event as a whole, and intersecting the Voronoi cell of the particle with a circle of radius R centred on the particle. Note that for the k_t algorithm (but not for Cambridge/Aachen or anti- k_t , nor in general for any other algorithm) the Voronoi area of a jet coincides with its passive area.

The area can be calculated either as a scalar, or as a 4-vector. Essentially the scalar case corresponds to counting the number of ghosts in the jet, while the 4-vector case corresponds to summing their

4-vectors (normalised such that for a narrow jet, the transverse component of the 4-vector is equal to the scalar area).

Jet areas are obtained by clustering with the class ClusterSequenceArea (rather than ClusterSequence, from which it is derived). Its constructor takes an AreaDefinition argument in addition to the list of particles and the JetDefinition.

It is worth noting that in the limit of very densely populated events, all area definitions tend to the same value [16]. It might therefore be advantageous to select a Voronoi area type, rather than an active one, when using areas for phenomenological tasks like pileup subtraction [17], as it generally requires less CPU time to calculate.

To summarise, in order to access the areas of the jets the user is exposed to two main classes:

```
class fastjet::AreaDefinition;
class fastjet::ClusterSequenceArea;
```

If jet areas are to be used to study the level of a diffuse noise which might be present in the event (like underlying event particles or pileup), a further specification, the phase space region over which to study such noise, will have to be given via the class

class fastjet::RangeDefinition;

These classes are described in detail below, and an example program is given in section 6.4.

6.1 fastjet::AreaDefinition

Area definitions are contained in fastjet::AreaDefinition class. Its two main constructors are:

for the various active and passive areas (which all involve ghosts) and

```
fastjet::AreaDefinition(fastjet::VoronoiAreaSpec voronoi_spec);
```

for the Voronoi area. A default constructor exists, and provides an active area with a ghost_spec that is suitable for a majority of area measurements with clustering algorithms and typical Tevatron and LHC rapidity coverage.

Information about the current AreaDefinition can be retrieved as follows:

```
/// return a description of the current area definition
std::string description() const ;
/// return info about the type of area being used by this defn
AreaType area_type() const ;
/// return a reference to the ghosted area spec (where relevant)
const GhostedAreaSpec & ghost_spec() const ;
/// return a reference to the voronoi area spec (where relevant)
const VoronoiAreaSpec & voronoi_spec() const ;
```

6.1.1 Ghosted Areas (active and passive)

There are two variants each of the active and passive areas, as defined by the AreaType enum:

The two active variants give identical results. The second one explicitly includes the ghosts when the user requests the constituents of a jet. The first of the passive variants explicitly runs through the procedure mentioned above, *i.e.* it clusters the events with one ghost at a time, and repeats this for very many ghosts. This can be quite slow, so we also provide the **passive_area** option, which makes use of information specific to the jet algorithm in order to speed up the passive-area determination.¹⁰

In order to carry out a clustering with a ghosted area determination, the user should also create an object that specifies how to distribute the ghosts.¹¹ This is done via the class fastjet::GhostedAreaSpec whose constructor is

```
fastjet::GhostedAreaSpec(double ghost_maxrap,
```

int	; repe	at	=	1,
dou	uble ghos	t_area	=	0.01,
dou	uble grid	_scatter	=	1.0,
dou	uble kt_s	catter	=	0.1,
dou	uble mean	_ghost_kt	=	1e-100);

The ghosts are distributed on a uniform grid in y and ϕ , with small random fluctuations to avoid degeneracies. The ghost_maxrap defines the maximum rapidity up to which ghosts are generated — typically jet areas will be reliable for jets up to rapidity $|y| \simeq \text{ghost_maxrap} - R$. The ghost_area is the area associated with a single ghost. The number of ghosts is inversely proportional to the ghost area, and so a smaller area leads to a longer CPU-time for clustering. However small ghost areas give more accurate results. We have found the default ghost area given above to be suitable for most applications.

For sparse events, the set of ghost particles that end up in a given jet is not unique, and depends on the degeneracy-breaking random shifts added to the ghost positions, as compared to a perfect grid distribution. To obtain a reliable area one may then repeat the area determination several times, the number of times being specified by the **repeat** variable. For hadron-level events a value of 5 is sufficient to give jet areas that are determined to within a few percent. In practice it is usually satisfactory even to set **repeat** = 1 and this is the default since it runs faster. For events with a dense distribution of true particles, there is no degeneracy in the ghost clustering and there is no need at all to use **repeat** > 1. If **repeat** > 1, a statistical uncertainty on the area, given by $\sigma/\sqrt{\text{repeat}-1}$, is provided for each jet. Note that the **repeat** value is ignored (*i.e.* taken to be 1) for **active_area_explicit_ghosts** and meaningless for the passive area in the k_t algorithm, which just calculates the Voronoi area discussed below (since they are identical).

¹⁰This ability is provided for k_t , Cambridge/Aachen, anti- k_t and the SISCone plugin. In the case of k_t it is actually a Voronoi area that is used, since this can be shown to be equivalent to the passive area [16]. For other algorithms it defaults back to the one_ghost_passive_area approach.

¹¹Or accept a default — which uses the default values listed in the explicit constructor and $ghost_maxrap = 6$

Other variables that the user may wish to set are: grid_scatter and kt_scatter, which are fractional random fluctuations of the position of the ghosts on the $y-\phi$ grid and of their transverse momentum; and mean_ghost_kt which is the average transverse momentum of the ghosts.

Even after the initialisation, the parameters can be modified by

```
void set_ghost_area (double);
void set_ghost_etamax (double);
void set_ghost_maxrap (double);
void set_grid_scatter (double);
void set_kt_scatter (double);
void set_mean_ghost_kt (double);
void set_repeat (int );
```

and information about the GhostedAreaSpec in use can be retrieved as follows:

```
/// for a summary
std::string description() const;
double ghost_etamax () const ;
double ghost_maxrap () const ;
double ghost_area () const ;
double grid_scatter () const ;
double kt_scatter () const ;
double mean_ghost_kt () const ;
int repeat () const ;
```

6.1.2 Voronoi Areas

The Voronoi areas of jets are evaluated by summing the corresponding Voronoi areas of the jets' constituents. The latter are obtained by considering the intersection between the Voronoi cell of each particle and a circle of radius R centred on the particle's position in the rapidity-azimuth plane.

The jets' Voronoi areas can be obtained from fastjet::ClusterSequenceArea by passing the proper fastjet::VoronoiAreaSpec specification to fastjet::AreaDefinition. Its constuctors are

```
/// default constructor (effective_Rfact = 1)
fastjet::VoronoiAreaSpec() ;
```

/// constructor that allows you to set effective_Rfact
fastjet::VoronoiAreaSpec(double effective_Rfact) ;

The second constructor allows one to modify (by a multiplicative factor effective_Rfact) the radius of the circle which is intersected with the Voronoi cells. With effective_Rfact = 1, for the k_t algorithm, the Voronoi area is equivalent to the passive area.

Information about the specification in use is returned by

```
/// return the value of effective_Rfact
double effective_Rfact() const ;
/// return a textual description of the area definition.
std::string description() const ;
```

The Voronoi areas are calculated with the help of Fortune's $(N \ln N)$ Voronoi diagram generator for planar static point sets [18].

6.2 fastjet::ClusterSequenceArea

This is the main class¹² to which the user is exposed for accessing cluster sequences that include information about jet areas. It is derived from fastjet::ClusterSequenceAreaBase (itself derived from fastjet::ClusterSequence) and includes the methods

```
/// return a reference to the area definition
virtual const fastjet::AreaDefinition & area_def() const ;
/// return the area associated with the given jet
virtual double area (const fastjet::PseudoJet & jet) const ;
/// return the error (uncertainty) associated with the determination
/// of the area of the jet; returns 0 when the repeat value = 1, and
/// also for the active_area_explicit_ghosts and certain passive areas
virtual double area_error (const fastjet::PseudoJet & jet) const ;
/// return a PseudoJet whose 4-vector is defined by the following integral
111
          \int dy d\phi PseudoJet(y, \phi, p_t = 1) * \Theta("y, \phi \text{ inside jet boundary"})
111
111
/// where PseudoJet(y, \phi, p_t = 1) is a 4-vector with the given
/// rapidity (y), azimuth (\phi) and p_t = 1, while \Theta("y, \phi inside jet boundary")
/// is a function that is 1 when y, \phi define a direction inside the
/// jet boundary and 0 otherwise.
111
virtual fastjet::PseudoJet area_4vector(const PseudoJet & jet) const ;
```

When the AreaType is active_area_explicit_ghosts, one may additionally use the following function

```
/// true if a jet is made exclusively of ghosts
virtual bool is_pure_ghost(const PseudoJet & jet) const;
```

to determine whether a jet is made purely of ghosts. Its argument can also be one of the constituents of a jet, in which case it will return **true** if that constituent is a ghost.

6.3 Areas and background subtraction

Jet areas can be used to study the level of a randomly distributed diffuse background which might be present together with the hard event of interest. After selecting a phase space region over which to analyse the jets, one can determine the average level rho (transverse momentum per unit area) of

¹² ClusterSequenceArea makes use of one among ClusterSequenceActiveArea, ClusterSequenceActiveAreaExplicitGhosts, ClusterSequencePassiveArea, ClusterSequence1GhostPassiveArea or ClusterSequenceVoronoiArea (all of them in the fastjet namespace of course), according to the choice made with AreaDefinition. The user might of course also use these classes directly.

the background, and also subtract it from the hard jets. This way of using jets to determine the noise level on an event-by-event basis was introduced and described in [17].

Note that we recommend that you use only the k_t or Cambridge/Aachen algorithms for calculating the UE/pileup density ρ . In fact, it is important to avoid algorithms with extreme behaviours for the areas. In particular one should not use anti- k_t (which has many jets with near-zero area) or SISCone (which can have jets with near-zero area, as well as monster jets with huge areas if the overlap parameter f is too small). In contrast the k_t and Cambridge/Aachen algorithms are suitable.

Once ρ has been determined using k_t or Cambridge/Aachen (and an appropriate value of R, typically of order 0.4–0.6, see [17] for more details), its value can then be used for UE/pileup subtraction in an analysis using any jet algorithm.

6.3.1 Fixed range

The fastjet::RangeDefinition class allows the user to set the rapidity-azimuth range over which he/she wishes to study the areas of the jets. It has two constructors. The first is

```
/// constructor for a range definition given by |y|<rapmax
fastjet::RangeDefinition(double rapmax);</pre>
```

which defines a range inclusive in azimuth $(0-2\pi)$, and bound by a maximum value for abs(y). The second constructor,

allows one to fully specify a rectangle in the rapidity-azimuth plane.

```
fastjet::RangeDefinition contains
```

```
/// return bool according to whether the jet is within the given range
bool is_in_range(const PseudoJet & jet) const ;
```

/// return bool according to whether the (rap,phi) point is within the given range
virtual bool is_in_range(double rap, double phi) const ;

/// (scalar) area of the range region
virtual double area() const ;

/// return the minimal and maximal rapidity of this range
virtual inline void get_rap_limits(double & rapmin, double & rapmax) const;

/// textual description of the range
virtual std::string description() const ;

6.3.2 Local Ranges

The virtual functions above can be overloaded by the user in a derived class, where a different range definition can be specified. An example can be found in CircularRange.hh, which implements a circular range centred on a jet's axis. Its constructors are

```
/// initialise CircularRange with a jet
fastjet::CircularRange(const fastjet::PseudoJet & jet, double distance) ;
/// initialise CircularRange with a (rap,phi) point
fastjet::CircularRange(double rap, double phi, double distance) ;
/// initialise CircularRange with just the radius parameter
fastjet::CircularRange(double distance) ;
```

This *local* range makes use of two further methods. First, it overrides a virtual function already present in RangeDefinition

```
/// For localizable classes override this function with a function
/// that returns true
virtual inline bool is_localizable() const ;
```

to return true. This causes the two following functions (present but disabled in the base class) to be enabled:

```
/// place the range on the rap-phi position
inline void set_position(const double & rap, const double & phi);
/// place the range on the jet position
inline void set_position(const PseudoJet & jet);
```

They allow one to place an existing local range at the supplied position.

Note: we expect the interface for ranges to evolve significantly in releases subsequent to 2.4.

6.3.3 Background subtraction

Once a range has been specified, the following methods, belonging to the base class ClusterSequenceAreaBase, are available for extracting the diffuse noise level and its fluctuations:

```
/// using jets within large (and with 4 vector areas if
/// use_area_4vector), calculate the median pt/area (\rho), as well as an
/// "error" (uncertainty), which is defined as the 1-sigma
/// half-width of the distribution of pt/A, obtained by looking for
/// the value \sigma such which a fraction (1-0.6827)/2 of the jets
/// (including empty jets) have p_t/A < \rho - \sigma \sqrt{\langle A \rangle}
```

```
111
/// The subtraction for a jet with uncorrected pt, p_t^U and area A is
111
    p_t^S = p_t^U - \rho A \pm \sigma \sqrt{A}
111
111
/// where the error is only that associated with the fluctuations
/// in the noise and not that associated with the noise having
/// caused changes in the hard-particle content of the jet.
111
/// NB: subtraction may also be done with 4-vector area of course,
/// and this is recommended for jets with larger values of R, as
/// long as rho has also been determined with a 4-vector area;
/// using a scalar area causes one to neglect terms of relative
/// order R^2/8 in the jet p_t.
void get_median_rho_and_sigma(const RangeDefinition & range,
                               bool use_area_4vector,
                               double & median, double & sigma,
                               double & mean_area);
```

A version exists also without the last argument.

A final version that is present allows one to calculate the median based on an explicit li:st of jets, rather than the cluster-sequence's inclusive_jets()

There are at least two ways this might be used. One is if you want to exclude (say) some number of hardest jets from the estimate of ρ :

```
// get list of jets
vector<PseudoJet> all_jets = sorted_by_pt(cs.inclusive_jets());
// remove the two hardest jets (you might do this more efficiently...)
all_jets.erase(all_jets.begin(), all_jets.begin()+2);
// Get rho based on all but the two hardest jets.
// The last argument MUST be set to true unless you are using explicit ghosts
cs.get_median_rho_and_sigma(all_jets, range, true, median, sigma, mean_area, true);
```

Another way it might be used is if you have the Cambridge/Aachen algorithm at some R value and wish to determine ρ based on the jets it would provide at a different jet radius, R' < R (without rerunning the clustering):

```
// get list of jets at smaller R' value
vector<PseudoJet> all_jets = cs.exclusive_jets(pow(R'/R,2)));
// The last argument MUST be set to false unless you are using explicit ghosts
cs.get_median_rho_and_sigma(all_jets, range, true, median, sigma, mean_area, false);
```

This is useful, for example, if you wish to determine ρ at a series of different angular resolutions R'. It only works for the Cambridge/Aachen algorithm. Note: the all_are_inclusive argument relates to FastJet's internal mechanism for dealing with empty area. The treatment of empty area is much more robust if the area is based on explicit ghosts (active_area_explicit_ghosts). Its use of strongly recommended for the above applications (in which case the value of all_are_inclusive becomes irrelevant).

There is also a routine for estimating rapidity-dependent densities, intended for example for heavyion events

This method is now considered *deprecated* (it is too sensitive to contamination from hard jets). An alternative is to use a *local* range definition, like the CircularRange mentioned above. Such a range definition will of course lead to a different median ρ according to where the range is placed. It is useful whenever the background distribution is expected to be non uniform in rapidity and, for instance in the case of non-central collisions, also in azimuth. With a wise choice for the distance *d*, CircularRange will then allow one to use ClusterSequenceArea::get_median_rho_and_sigma() on a jet-by-jet basis. One can similarly also use a plain RangeDefinition in a limited rapidity and/or azimuth region.

Note that more specialised methods for extracting the noise level might be present at the level of a specific class, e.g. ClusterSequenceActiveArea. At least some of them should however be considered obsolete, and might be removed in future releases of FastJet.

Finally, the base class ClusterSequenceAreaBase contains virtual methods (usually overloaded by the derived classes) that deal with pure-ghost jets that might be formed when adding ghost particles to the events in order to calculate ghosted areas:

```
/// return the total area, within range, that is free of jets
virtual double empty_area(const RangeDefinition & range) const;
/// return something similar to the number of pure ghost jets
/// in the given range in an active area case.
/// Note that the number returned is a double.
virtual double n_empty_jets(const RangeDefinition & range) const;
```

With active area definitions these are calculated based on the observed number of pure ghost jets (and unclustered ghosts) in range; for Voronoi and passive and areas, they are calculated using the difference between the total range area and the area of the jets contained in the range, with the number of empty jets then being calculated based on the average jet area.

6.4 Example program with areas and subtraction

The following simplified example program clusters a series of input particles into jets, evaluates the median transverse momentum per unit area of the event (both for the standard area, and the alternative 4-vector area), and performs the subtraction of the background noise from the hard jets.

```
#include "fastjet/ClusterSequenceArea.hh"
namespace fj = fastjet;
using namespace std;
int main (int argc, char ** argv) {
  // select the jet algorithm (kt with R=0.7)
  double R = 0.7;
  fj::JetDefinition jet_def(fj::kt_algorithm,R);
  // select the area type (active area with default ghosts)
  fj::AreaDefinition area_def(fj::active_area);
  // read in input particles
  double px, py , pz, E;
  vector<fj::PseudoJet> input_particles;
  while (cin >> px >> py >> pz >> E) {
    input_particles.push_back(fj::PseudoJet(px,py,pz,E));
  }
  // cluster the event, also obtaining area information
  fj::ClusterSequenceArea csa(input_particles,jet_def,area_def);
  // get list of jets with pt > 5 GeV
  vector<fj::PseudoJet> jets = csa.inclusive_jets(5.0);
  // set the rapidity-azimuth range within which to study background
  double rapmax = 4.0;
  fj::RangeDefinition range(rapmax);
  // get median pt of all jets using standard area
  double median_pt = csa.median_pt_per_unit_area(range);
  // get median pt of all jets using area_4vector
  double median_pt_4vect = csa.median_pt_per_unit_area_4vector(range);
  cout << "\n Median rho (pt per unit area) = " << median_pt << " GeV\n\n";</pre>
                                          area pt_sub pt_sub_4vect " << endl;</pre>
  cout << " i
                  rap
                          phi
                                    pt
  // perform subtraction, output results
    for (unsigned i = 0; i < jets.size(); i++) {</pre>
       // get area of jet i
       double area = csa.area(jets[i]);
       // get area_4vector of jet i
       fj::PseudoJet area4vect = csa.area_4vector(jets[i]);
       // standard subtraction
       double pt_sub = jets[i].perp() - median_pt*area;
       // 4-vector subtraction
       fj::PseudoJet jet_sub = jets[i] - median_pt_4vect*area4vect;
       // print out results
```

Note that the above listing ignores the issue of cases where the transverse momentum to be subtracted is larger than the jet's p_t . This occurs when the jet is not really a true hard jet, but rather mostly composed of the background contamination. In such cases, the subtraction should not be performed, and the jet should be instead be discarded.

For convenience, the above operations are pre-packaged together with a certain number of safety checks (in particular if the subtracted transverse momentum is too large, routines that return a jet, set all of its components to zero. Methods that are given a RangeDefinition object in input, and therefore calculate ρ themselves, do it according to area_4vector, unless a further input flag is present and false):

```
/// return a vector of all subtracted jets, using area_4vector, given rho.
/// Only inclusive_jets above ptmin are subtracted and returned.
/// The ordering is the same as that of sorted_by_pt(cs.inclusive_jets()),
/// i.e. not necessarily ordered in pt once subtracted
std::vector<PseudoJet> subtracted_jets(const double rho,
                                       const double ptmin=0.0) const;
/// return a vector of subtracted jets, using area_4vector.
/// Only inclusive_jets above ptmin are subtracted and returned.
/// The ordering is the same as that of sorted_by_pt(cs.inclusive_jets()),
/// i.e. not necessarily ordered in pt once subtracted
std::vector<PseudoJet> subtracted_jets(const RangeDefinition & range,
                                       const double ptmin=0.0) const;
/// return a subtracted jet, using area_4vector, given rho
PseudoJet subtracted_jet(const PseudoJet & jet, const double rho) const;
/// return a subtracted jet, using area_4vector; note
/// that this is potentially inefficient if repeatedly used for many
/// different jets, because rho will be recalculated each time around.
PseudoJet subtracted_jet(const PseudoJet & jet,
                         const RangeDefinition & range) const;
/// return the subtracted pt, given rho
double subtracted_pt(const PseudoJet & jet, const double rho,
                     bool use_area_4vector=false) const;
/// return the subtracted pt; note that this is
/// potentially inefficient if repeatedly used for many different
/// jets, because rho will be recalculated each time around.
```

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7 Plugin jet algorithms

It can be useful to have a common interface for a range of jet algorithms beyond the native $(k_t, \text{anti-}k_t \text{ and Cambridge/Aachen})$ algorithms, and it can also be useful to use the area-measurement tools for these other jet algorithms. In order to facilitate this, the FastJet package provides a *plugin* facility, allowing almost any other jet algorithm¹³ to be used within the same framework.

7.1 Generic plugin use and construction

Any plugin is to be derived from the abstract base class fastjet::JetDefinition::Plugin,

```
class JetDefinition::Plugin{
public:
  /// return a textual description of the jet-definition implemented
  /// in this plugin
  virtual std::string description() const = 0;
  /// given a ClusterSequence that has been filled up with initial
  /// particles, the following function should fill up the rest of the
  /// ClusterSequence, using the following member functions of
  /// ClusterSequence:
       - plugin_do_ij_recombination(...)
  ///
  ///
       - plugin_do_iB_recombination(...)
  virtual void run_clustering(ClusterSequence &) const = 0;
  /// a destructor to be replaced if necessary in derived classes...
 virtual ~Plugin() {};
 //----- ignore what follows for simple usage! ------
  /// return true if there is passive areas can be efficiently determined by
  /// (a) setting the ghost_separation scale (see below)
  /// (b) clustering with many ghosts with p_t \ll ghost_separation_scale
  /// (c) counting how many ghosts end up in a given jet
 virtual bool supports_ghosted_passive_areas() const {return false;}
 /// set the ghost separation scale for passive area determinations
  /// in future runs (NB: const, so should set internal mutable var)
 virtual void set_ghost_separation_scale(double scale) const;
 virtual double ghost_separation_scale() const;
```

};

A JetDefinition can be constructed by providing a pointer to a JetDefinition::Plugin; the resulting JetDefinition object can then be used identically to the normal JetDefinition objects used in the previous sections.

// have some plugin class derived from the Plugin base class
class CDFMidPointPlugin : public fastjet::JetDefinition::Plugin {...};

¹³Except those that perform $3 \rightarrow 2$ clusterings for which there is no unique mapping of particles into jets (some particles are effectively shared among more than one jet).

```
// create an instance of the CDFMidPointPlugin class
CDFMidPointPlugin cdf_midpoint( [... options ...] );
// create the jet definition
fastjet::JetDefinition jet_def = fastjet::JetDefinition( & cdf_midpoint);
// then create ClusterSequence with the input particles and jet_def,
// and use it to extract jets as usual
```

Any plugin class must define the description and run_clustering member functions. The former just returns a textual description of the jet algorithm and its options (e.g. radius, etc.), while the latter does the hard work of running the user's own jet algorithm and transferring the information to the ClusterSequence class. This is best illustrated with an example:

using namespace fastjet;

```
void CDFMidPointPlugin::run_clustering(ClusterSequence & clust_seq) {
    // when run_clustering is called, the clust_seq has already been
    // filled with the initial particles, which are available through the
    // jets() array
    const vector<PseudoJet> & initial_particles = clust_seq.jets();
    // it is up to the user to do their own clustering on these initial
    // particles
```

```
// ...
```

Once the plugin has run its own clustering it must transfer the information back to the clust_seq. This is done by recording mergings between pairs of particles or between a particle and the beam. The new momenta are stored in the clust_seq.jets() vector, after the initial particles. Note though that the plugin is not allowed to modify clust_seq.jets() itself. Instead it must tell clust_seq what recombinations have occurred, via the following (ClusterSequence member) functions

/// jets()[jet_i] and the beam, with the specified diB; when looking

```
/// for inclusive jets, any iB recombination will returned to the user
/// as a jet.
void plugin_record_iB_recombination(int jet_i, double diB);
```

These dij recombination functions return the index newjet_k of the newly formed pseudojet. The plugin may need to keep track of this index in order to specify subsequent recombinations.

Certain (cone) jet algorithms do not perform pairwise clustering — in these cases the plugin must invent a ficititious series of pairwise recombinations that leads to the same final jets. Such jet algorithms may also produce extra information that cannot be encoded in this way (for example a list of stable cones), but to which one may still want access. For this purpose, during run_clustering(...), the plugin may call the ClusterSequence member function:

inline void plugin_associate_extras(std::auto_ptr<ClusterSequence::Extras> extras);

where ClusterSequence::Extras is a dummy class which the plugin should extend so as to provide the relevant information:

```
class ClusterSequence::Extras {
  public:
    virtual ~Extras() {}
    virtual std::string description() const;
};
```

A method of ClusterSequence then provides the user with access to the extra information:

```
/// returns a pointer to the extras object (may be null) const
Extras * extras() const;
```

The user should carry out a dynamic cast so as to convert the extras back to the specific plugin extras class, as illustrated later for SISCone.

Example plugin jet algorithms that are provided with FastJet are the CDFMidPointPlugin illustrated above and also a CDFJetCluPlugin. These interface code available at [19]. The PxConePlugin interfaces the PxCone code [20]. All three of these cone jet algorithms, though widely used, are infrared or collinear unsafe (depending on the parameters). The SISConePlugin, described in detail below, interfaces the infrared safe seedless cone algorithm of [7]. Examples using these plugins and some further documentation are provided in the plugins/ directory.

7.2 SISCone Plugin

SISCone [7] is an implementation of a cone type jet algorithm. As with most modern cone algorithms, it is divided into two parts: first it searches for stable cones; then, because a particle can appear in more than one stable cone, a 'split-merge' procedure is applied, which ensures that no particle ends up in more than one jet. The stable cones are identified using an $\mathcal{O}(N^2 \ln N)$ seedless approach. This (and some care in the the 'split-merge' procedure) ensures that the jets it produces are insensitive to additional soft particles and collinear splittings (i.e. the jets are infrared and collinear safe).

The plugin library and include files are to be be found in the plugins/SISCone directory, and the main header file is SISConePlugin.hh. The SISConePlugin class has a constructor with the following structure

```
SISConePlugin (double cone_radius,
```

A cone centered at y_c , ϕ_c is stable if the sum of momenta of all particles *i* satisfying $\Delta y_{ic}^2 + \Delta \phi_{ic}^2 < \text{cone_radius}^2$ has rapidity y_c , ϕ_c . The overlap_threshold is the fraction of overlapping momentum above which two protojets are merged in a Tevatron Run II type [8] split-merge procedure.¹⁴ The radius and overlap parameters are a common feature of most modern cone algorithms. Because some event particles are not to be found in any stable cone [21], SISCone can carry out multiple stable-cone search passes (as advocated in [22]): in each pass one searches for stable cones using just the subset of particles not present in any stable cone in earlier passes. Up to n_pass_max passes are carried out, and the algorithm automatically stops at the highest pass that gives no new stable cones. The default of n_pass_max = 0 is equivalent to setting it to ∞ . Since concern has been expressed that an excessive number of stable cones may complicate cone jets in events with high noise [8], the protojet_ptmin parameter allows one to use only protojets with $p_t \geq \text{protojet_ptmin}$ in the split-merge phase (all others are thrown away).¹⁵

In many cases SISCone's most time-consuming step is the search for stable cones. If one has multiple SISConePlugin-based jet definitions, each with caching=true, a check will be carried out whether the previously clustered event had the same set of particles and the same cone radius and number of passes. If it did, the stable cones are simply reused from the previous event, rather than being recalculated, and only the split-merge step is repeated, often leading to considerable speed gains.

A final comment concerns the split_merge_scale parameter. This controls both the scale used for ordering the protojets during the split-merge step during the split-merge step, and also the scale used to measure the degree of overlap between protojets. While various options have been implemented,

enum SplitMergeScale {SM_pt, SM_Et, SM_mt, SM_pttilde };

we recommend using only the last of them $\tilde{p}_t = \sum_{i \in jet} |p_{t,i}|$, which is also the default scale. The other scales are included only for historical comparison purposes: p_t (used in several other codes) is IR unsafe for events whose hadronic component conserves momentum, E_t (advocated in [8]) is not boost-invariant, and $m_t = \sqrt{m^2 + p_t^2}$ is IR unsafe for events whose hadronic component conserves momentum and stems from the decay of two identical particles.

An example of the use of the SISCone plugin would be as follows:

// define a SISCone plugin pointer
fastjet::SISConePlugin * plugin;

```
// allocate a new plugin for SISCone
double cone_radius = 0.7;
```

¹⁴Though its default value is 0.5 (retained for backwards compatibility of the interface) we strongly recommend using a higher value, e.g. 0.75, especially in high-noise environments, in order to disfavour the production of monster jets through repeated merge operations.

¹⁵Early experience indicates that protojet_ptmin is actually perfectly adequate and that potential problems of massively agglomerated jets that can occur in high-noise environments (for a wide range of cone algorithms) can be addressed with a slightly larger value of the overlap_threshold, ≥ 0.6 .

```
double overlap_threshold = 0.5;
plugin = new fastjet::SISConePlugin (cone_radius, overlap_threshold);
// create a jet-definition based on the plugin
fastjet::JetDefinition jet_def(plugin);
// prepare the set of particles
vector<fastjet::PseudoJet> particles;
read_input_particles(cin, particles); // or whatever you want here
// run the jet algorithm and look at the jets
fastjet::ClusterSequence clust_seq(particles, jet_def);
vector<fastjet::PseudoJet> inclusive_jets = clust_seq.inclusive_jets();
// then analyse the jets as for native FastJet jets
// only when you will no longer be using the jet definition, or
// ClusterSequence objects that involve it, may you delete the
// plugin
```

```
delete plugin;
```

Note that the it makes no sense to ask for exclusive jets from a SISCone based ClusterSequence.

Some extra output information is appropriate for a cone algorithm that is not of relevance in clustering algorithms, through the extras resource,

To determine the pass at which a given jet was found, one does the following

```
int pass = extras->pass(jet);
```

The user may also obtain a list of the positions of original stable cones as follows:

```
vector<fastjet::PseudoJet> stable_cones(extras->stable_cones());
```

The stable cones are represented as four-momenta, for which only the rapidity and azimuth are meaningful. The user_index() indicates the pass at which a given stable cone was found.

In the current version of SISCone, the user_index() of a jet also corresponds to the pass at which it was found, however this manner of accessing the pass for a jet is *deprecated* (for reasons related to the internal representation of jets, it fails for single-particle jets). It is retained in version 2.4 for backwards compatibility, but will be removed at some stage in the future.

SISCone uses *E*-scheme recombination internally and also for constructing the final jets from the list of constituents. For the latter task, the user may instead instruct SISCone to use the jet-definition's own recombiner, with the command

```
plugin->set_use_jet_def_recombiner(true);
```

In this case the user_index() no longer contains the information about the pass.

Since SISCone is infrared safe, it may meaningfully be used also with the ClusterSequenceArea class. Note however that in that case ones loses the cone-specific information from the jets, because of the way FastJet filters out the information relating to ghosts in the clustering. If the user needs both areas and cone-specific information, she/he may use the

ClusterSequenceActiveAreaExplicitGhosts class (for usage information, see the corresponding .hh file).

A final remark concerns speed and memory requirements: as mentioned above, SISCone takes $\mathcal{O}(N^2 \ln N)$ time to find jets, and the memory use is $\mathcal{O}(N^2)$; taking $N = 10^3$ as a reference point, it runs in a few tenths of a second, making it about 100 times slower than native FastJet algorithms. These are 'expected' results, i.e. valid for a suitably random set of particles. In area determinations, the ghost particles are anything but random, and run times and memory usage are, in practice, somewhat larger than for a normal QCD event with the same number of particles. We therefore recommend running with not too small a ghost_area (e.g. ~ 0.05) and using grid_scatter = 1, which helps to reduce the number of stable cones (and correspondingly, the time and memory usage of the subsequent split-merge step). An alternative, which has been found to be acceptable in most situations, is to use a passive area, since this is relatively fast to calculate with SISCone.

7.3 Other plugins for *pp*

Not all plugins are enabled by default in FastJet. At configuration time ./configure.sh --help will indicate which ones get enabled by default. To enable all plugins, run configure with the argument --enable-allplugins, while to enable all but PxCone (which requires fortran, and can introduce link-time issues) use --enable-allcxxplugins.

All plugins are in the fastjet namespace. Below we show the file that needs to be included and the constructor for each plugin.

Except where stated, the usual way to access jets from these plugins is through ClusterSequence::inclusive_jets().

Most of the algorithms listed below are either infrared (IR) or collinear unsafe. The details are indicated for each algorithm as follows: IR_{n+1} means that the hard jets may be modified if, to an ensemble of n hard particles in a common neighbourhood, one adds a single soft particle; $Coll_{n+1}$ means that for n hard particles in a common neighbourhood, the collinear splitting of one of them may modify the hard jets. The FastJet authors (and numerous theory-experiment accords) advise against the use IR and collinear safe jet algorithms. Interfaces to these algorithms have been provided mainly for legacy comparison purposes.

As of FastJet version 2.4, this section is partially incomplete (in particular it misses many references). This will hopefully evolve for future versions.

7.3.1 CDF Midpoint.

One of the two algorithms used by CDF in Run II of the Tevatron, based on [8]. It is a midpoint-type iterative cone with a split-merge step.

The overlap threshold (f) used by CDF is usually 0.5, the seed threshold is 1 GeV and in most measurements the cone area fraction is 1. With an area fraction < 1 this becomes the searchcone

algorithm of [21].

Further control over the plugin can be obtained by consulting the header file.

The underlying code for this algorithm was taken from a webpage provided by Joey Huston (with minor modifications to ensure reasonable run times with optimising compilers for 32-bit intel processors — these modifications do not affect the final jets).

Note: this algorithm is IR_{3+1} unsafe (in the limit of zero seed threshold [7]; with cone_area_fraction \neq 1 it becomes IR_{2+1} unsafe [22]). It is to be deprecated for new experimental or theoretical analyses.

7.3.2 CDF JetClu.

The other algorithm used by CDF during Run II, as well as their main algorithm during Run I.

This is an iterative cone with split-merge and optional "ratcheting" if iratch == 1 (particles that appear in one iteration fo a cone are retained in future iterations). The overlap threshold is usually set to 0.75 in CDF analyses.

Further control over the plugin can be obtained by consulting the header file.

The underlying code for this algorithm was taken from a webpage provided by Joey Huston.

Note: this algorithm is IR_{2+1} unsafe (and some IR unsafety persists with non-zero seed threshold). It is to be deprecated for new experimental or theoretical analyses. Note also that the underlying implementation groups particles ogether into calorimeter towers (with CDF-type geometry) before running the jet algorithm.

7.3.3 DØ Run II cone

The main algorithm used by $D\emptyset$ in Run II of the Tevatron, which is a midpoint type iterative cone with split-merge.

Instead of a seed threshold, the algorithm places a cut on the minimum E_t of the cones during iteration (related to min_jet_Et). The split_ratio is the same as the overlap threshold in other split-merge based algorithms (DØ usually use 0.5). It is the FastJet authors' understanding that two values have been used for min_jet_Et, 8 GeV (in earlier publications) and 6 GeV (in more recent publications).

The underlying code for this algorithm was provided by Lars Sonnenschein.

Note: this algorithm is IR_{3+1} unsafe (IR_{2+1} for jets with energy too close to min_jet_Et). It is to be deprecated for new experimental or theoretical analyses.

7.3.4 ATLAS iterative cone

The (iterative) cone (with split-merge) algorithm used by ATLAS during the preparation for the LHC.

f is the overlap threshold

The underlying code for this algorithm was extracted from SpartyJet [27].

Note: this algorithm is IR_{2+1} unsafe (in the limit of zero seed threshold). It is to be deprecated for new experimental or theoretical analyses.

7.3.5 CMS iterative cone

The (iterative) cone (with progressive removal) algorithm used by CMS during the preparation for the LHC.

```
#include ''fastjet/CMSIterativeConePlugin.hh''
///...
CMSIterativeConePlugin (double ConeRadius, double SeedThreshold=0.0);
```

The underlying code for this algorithm was extracted from the CMSSW web site, with certain small service routines having been rewritten by the FastJet authors. The resulting code was validated by clustering 1000 events with the original version of the CMS software and comparing the output to the clustering performed with the FastJet plugin. The jet contents were identical in all cases. However the jet momenta differed at a relative precision level of 10^{-7} , related to the use of single-precision arithmetic at some internal stage of the CMS software (while the FastJet version is in double precision).

Note: this algorithm is Coll_{3+1} unsafe [6]. It is to be deprecated for new experimental or theoretical analyses.

7.3.6 PxCone

A fortran plugin for the PxCone algorithm, which is an iterative cone with midpoints and a split-drop procedure

with a threshold on the minimum cone transverse energy if it is to be included in the split-drop stage. If E_scheme_jets is true then the plugin applies an E-scheme recombination to provide the momenta of the final jets (by default an E_t type recombination scheme is used).

The base code for this plugin is written in Fortran and, on some systems, problems have been reported at the link stage due to mixing Fortran and C++. The Fortran code has been modified by the FastJet authors to provide the same jets regardless of the order of the input particles. This involved a small modification of the midpoint procedure, which can have a minor effect on the final jets and should make the algorithm correspond to the description of [24].

The underlying code for this algorithm was taken from a google search for PxCone!

Note: this algorithm is IR_{3+1} unsafe. It is to be deprecated for new experimental or theoretical analyses.

7.3.7 TrackJet

This algorithm has been used at the Tevatron for identifying jets from charged-particle tracks in underlying-event studies (a citation is needed here!).

Two recombination schemes are involved: the first one indicates how momenta are recombined to provide the final jets (once particle-jet assignments are known), the second one indicates how momenta are combined in the procedure that constructs the jets.

The underlying code for this algorithm was written by the FastJet authors, based on code extracts from the Rivet implementation, written by Andy Buckley with input from Manuel Bahr and Rick Field.

Note: this algorithm is believed to be $Coll_{3+1}$ unsafe. It is to be deprecated for new experimental or theoretical analyses.

7.4 Other plugins for e^+e^-

7.4.1 Cambridge algorithm

The original e^+e^- cambridge [4] algorithm is provided as a plugin:

```
#include ''fastjet/EECambridgePlugin.hh''
///...
EECambridgePlugin (double ycut);
```

This algorithms performs sequential recombination of the pair of particles that is closest in angle, except when $y_{ij} = \frac{2\min(E_i^2, E_j^2)}{Q^2}(1 - \cos \theta) > y_{cut}$, in which case the less energetic of *i* and *j* is labelled a jet, and the other member of the pair remains free to cluster.

To access the jets, the user should use the $inclusive_jets()$, *i.e.* as they would for the majority of the pp algorithms.

The underlying code for this algorithm was written by the FastJet authors.

7.4.2 Jade algorithm

The JADE algorithm [25, 26], a sequential recombination algorithm with distance measure $d_{ij} = 2E_i E_j (1 - \cos \theta)$, is available through

```
#include ''fastjet/JadePlugin.hh''
///...
JadePlugin ();
```

To access the jets at a given $y_{cut} = d_{cut}/Q^2$, the user should call ClusterSequence::exclusive_jets_ycut(double ycut).

Note: the JADE algorithm has been used with various recombination schemes. The current plugin will use whatever recombination scheme the user specifies with for the jet definition. The default *E*-scheme is what was used in the original JADE publication [25]. To modify the recombination scheme, the user may first construct the jet definition and then use either of

void JetDefinition::set_recombination_scheme(RecombinationScheme recomb_scheme); void JetDefinition::set_recombiner(const Recombiner * recomb)

(cf. sections 5.6,A) to modify the recombination scheme.

The underlying code for this algorithm was written by the FastJet authors.

7.5 Building new sequential recombination algorithms

To enable users to more easily build plugins for new sequential recombination algorithms, fastjet also provides a class NNH, which provides users with access to an implementation of the nearest-neighbour heuristic for establishing and maintaining information about the closest pair of objects in a dynamic set of objects (see [23] for an introduction to this and other generic algorithms). In good cases this allows one to construct clustering that runs in N^2 time, though its worst case can be as bad as N^3 . It is a templated class and the template argument should be a class that stores the minimal information for each jet so as to be able to calculate interjet distances. It underlies the implementations of the Jade and e^+e^- Cambridge plugins. The interested user should consult those codes for more information, as well as the header for the NNH class.

8 Compilation notes

Compilation and installation make use of the standard

```
% ./configure
```

```
% make
```

```
\% make check
```

```
% make install
```

procedure. Explanations of available options are given in the INSTALL file in the top directory.

In order to access the NlnN strategy for the k_t algorithm the library needs to be compiled with the Computational Geometry Algorithms Library CGAL [15].¹⁶ At configure time the --enable-cgal

¹⁶This same strategy gives $N \ln N$ performance for Cambridge/Aachen and $N^{3/2}$ performance for anti- k_t (whose sequence for jet clustering triggers a worst-case scenario for the underlying computational geometry methods.)

option may be used to specify that CGAL support should be included.

CGAL may be obtained in source form from http://www.cgal.org/. Under linux, with CGAL versions 3.2 and 3.3, after compilation and installation, the user will be encouraged to set an environment variable CGAL_MAKEFILE, which points to the Makefile generated by CGAL at install time, which contains various definitions of locations of include files. The user may specify the location of this file to FastJet either through the above environment variable, or with the --with-cgalmakefile=... configuration option. For CGAL 3.4 the user should instead specify --with-cgaldir=... unless the CGAL files are installed in a standard location.

The NlnNCam strategy does not require CGAL, since it is based on a considerably simpler computational-geometry structure [12].

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A External Recombination Schemes

If the user wishes to introduce a new recombination scheme, she may do so writing a class derived from JetDefinition::Recombiner:

A jet definition can then be constructed by providing a pointer to an object derived from JetDefinition::Recombiner instead of the RecombinationScheme index:

The derived class JetDefinition::DefaultRecombiner is what is used internally to implement the various recombination schemes if an external Recombiner is not provided. It provides a useful example of how to implement a new Recombiner class.

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