GPU Acceleration and EDM Developments for the ATLAS 3D Calorimeter Clustering in the Software Trigger

CHEP 2024

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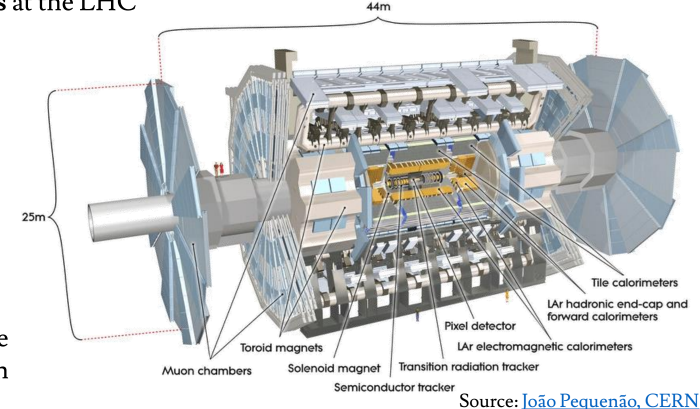




Context

The ATLAS Experiment

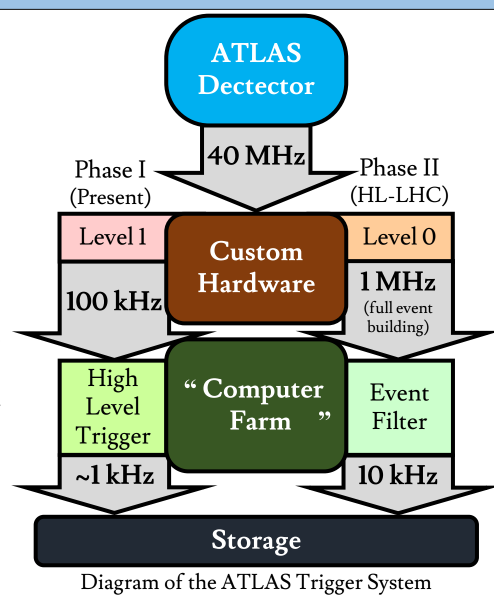
- <u>**A** T</u>oroidal <u>**L**HC <u>**A**</u>pparatu<u>s</u>[1]</u>
- One of the two **general-purpose detectors** at the LHC
- Three layers:
 - Inner Detector
 - Calorimeters
 - Muon Spectrometers
- 10⁸ electronic channels
 - 187652 calorimeter cells with multiple gain paths to optimize resolution *versus* dynamic range of operation



[1] - The ATLAS Experiment at the CERN Large Hadron Collider, DOI 10.1088/1748-0221/3/08/S08003

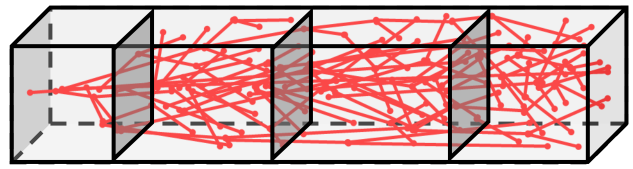
The ATLAS Trigger

- The ATLAS Trigger^[1] is used to filter the detected events to ensure a manageable output rate
 - Speed *versus* accuracy trade-offs can be relevant
- Two stages:
 - Hardware-based (Level 1/Level 0)
 - Software-based (High-Level Trigger/Event Filter)
- The High-Luminosity LHC Upgrade will increase the luminosity, making event reconstruction more computationally demanding
- The Phase II upgrade needed for the High-Luminosity LHC increases event rate at the software-based stage by a factor of 10
- This **higher computational load** requires **more computing power** available for the trigger and/or **better optimization**
- Alternative: hardware acceleration
 - Ongoing studies for both FPGA and GPU acceleration
- [1] The ATLAS trigger system for LHC Run 3 and trigger performance in 2022, DOI 10.1088/1748-0221/19/06/P06029

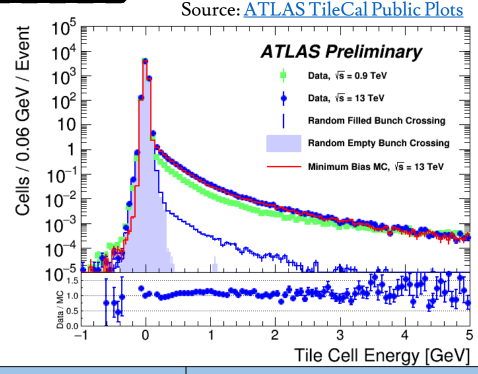


Calorimeter Reconstruction Algorithms

• Reconstruction of **showers** generated by outgoing particles in the calorimeters of the ATLAS experiment

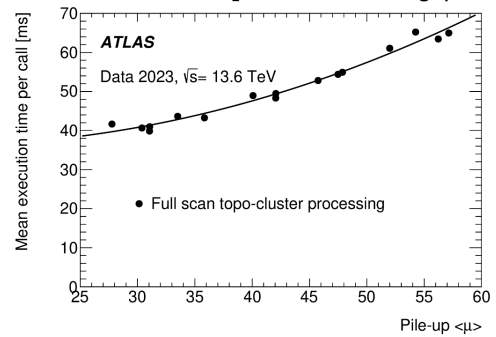


- Showers deposit their energy in a finite region of space: a calorimeter cell
- Calorimeter cells organized in up to 28 sampling layers/regions
- Two main sources of **noise**: electronic read-out and pile-up
 - The **noise estimate** is typically a function of the gain of the cell
 - For the **Tile calorimeter**, the electronic noise can be estimated by a **two-Gaussian model**, which involves more sophisticated computations (inverse error function of error functions)



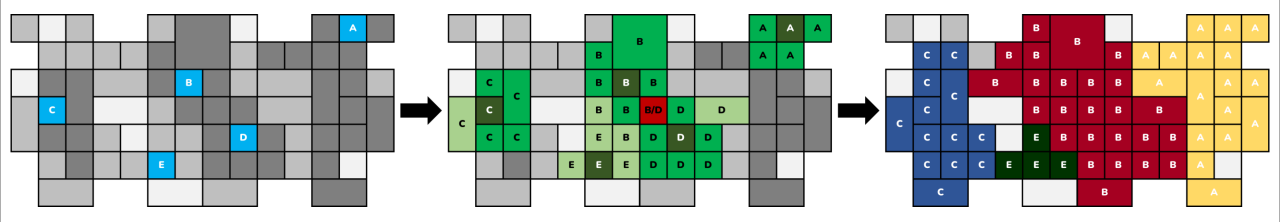
Topological and Topo-Automaton Clustering

- Topological Clustering is the currently used approach for calorimeter reconstruction in ATLAS
- Main criterion for assigning cells to the clusters is the signal-to-noise ratio (SNR) of the energy deposition
 - This is essentially the **relevance** of the **contribution** of each cell to the **reconstruction of the underlying physics**
- Clustering typically groups up several tens of cells, some clusters may be significantly larger
- Several hundred to a few thousand clusters per event, depending on the physical process
- Significant dependence on the number of collisions per bunch crossing (μ) in terms of the execution time

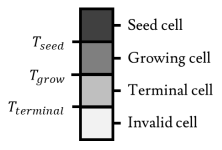


Source: The ATLAS trigger system for LHC Run 3 and trigger performance in 2022, DOI 10.1088/1748-0221/19/06/P06029

- Two main algorithmic stages:
 - Cluster growing: iteratively assign cells to clusters based on the SNR

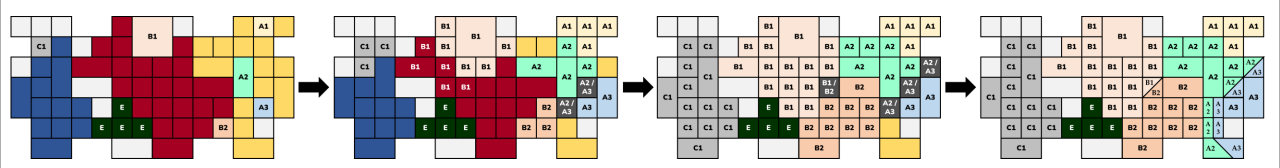


> Classify cells as **seed**, **growing** or **terminal**



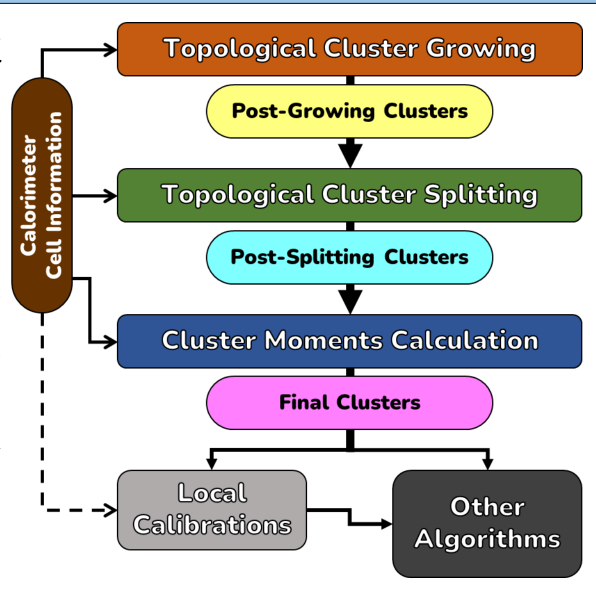
- > Clusters grow out from the seeds to their neighbouring cells in an order defined by the SNR of the seed
- > Clusters are merged if they touch through growing cells

- Two main algorithmic stages:
 - Cluster splitting: split the clusters around local maxima of the energy to distinguish different objects travelling in the same direction



- > Identify local maxima
- Exclude maxima from certain regions of the detector that overlap in certain directions to favour layers with greater radiation depth
- > Start growing the clusters to neighbouring cells in an order defined by the energy of the cells
- > Cells that can belong to more than one maximum are shared
- > Shared cells grow clusters only in the end and are weighted based on the energy and distance to the centroid: $w_1 = \frac{E_1}{E_1 + r E_2}$, $w_2 = 1 w_1$, with $r = e^{d_1 d_2}$

- A final step involves calculating several cluster moments, based on weighted sums of the properties of the cells and functions thereof
 - Some of the moments require calculating eigenvalues and eigenvectors of a 3 × 3 matrix
- Some **local calibrations** may be applied to the clusters, based on the cluster moments and the constituent cells
- Topological Clustering is the most computationally demanding algorithm of the calorimeter reconstruction and among the top 20th most computationally demanding algorithms within the ATLAS trigger
 - The same implementation is used for both online and offline reconstruction (with potentially different configuration)
- Topological clustering, as described, is not acceleratorfriendly: a different algorithmic approach is needed



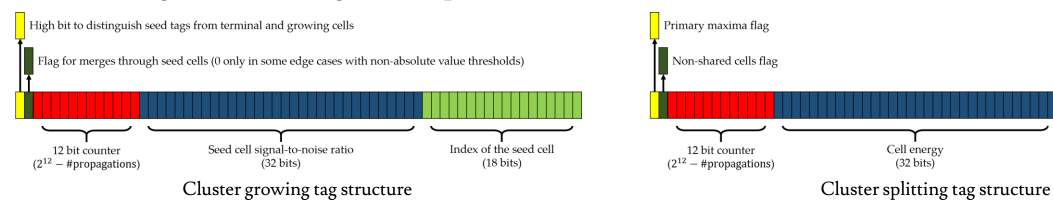
Limitations of Topological Clustering

- The clusters are expressed as lists of cells¹ which must be resized as they grow
- The algorithm itself involves keeping track of multiple lists of cells¹, especially for cluster splitting
- Resizing a container is difficult to do in parallel, and it goes against the memory model of both GPUs and FPGAs
- For a more parallel-friendly implementation, we can instead mark the cells that belong to each cluster with a "tag"
 - By constructing these tags appropriately, the sorting steps can be skipped entirely: floating point numbers that follow the IEEE-754 standard can be put in a "total ordering" where the bit patterns, interpreted as integers, are ordered in the same way as the original floating point numbers
 - By defining a set of rules for how these tags are propagated from a cell to its neighbours, one can replicate the entire behaviour of the iterative parts of cluster growing and cluster splitting while only considering each pair of neighbours independently from each other (potentially in parallel, as long as tag updates are thread-safe)
- Since we have both a **state** for each cell and can specify the **rules for how that state changes** based on the neighbourhood, this is equivalent to a **cellular automaton**, hence **Topo-Automaton Clustering**

^{1-&}quot;List" is used here in the sense of an ordered collection of items; specifically, they correspond to dynamically allocated arrays, or "vectors" in C++.

Topo-Automaton Clustering

• Cluster tags are 64-bit integers with specific structure:



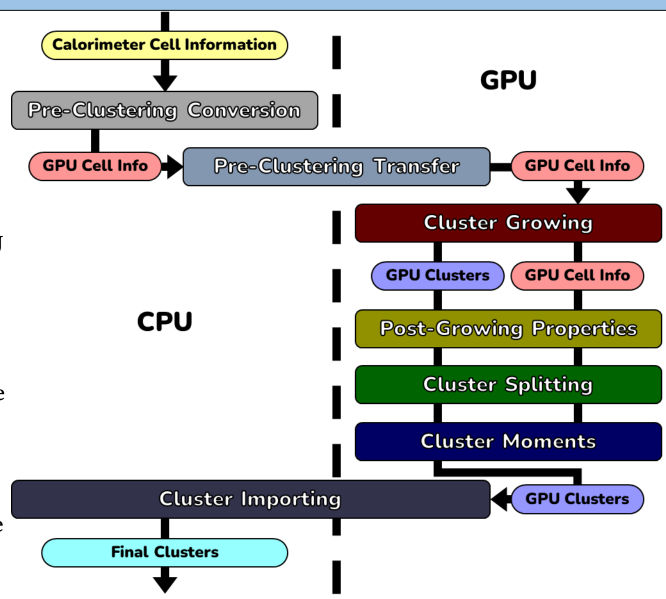
- The tags are propagated through pairs of neighbouring cells satisfying the conditions for clusters to expand
 - We handle each pair of cells in parallel, using appropriate atomic operations when needed
- Additional logic (e. g. keeping a cell to cluster index table) reduces the number of iterations
- All necessary temporary information stored in the same block of memory meant to hold the cluster moments
 (calculated only at the end), everything can be pre-allocated
 - Total **per event memory** footprint is ~**80 MB** (per CPU thread)
 - Cell geometry, neighbourhood relations and noise constants represent ~100 MB of constant information

Index of the cell

(or cluster for non-split clusters) (18 bits)

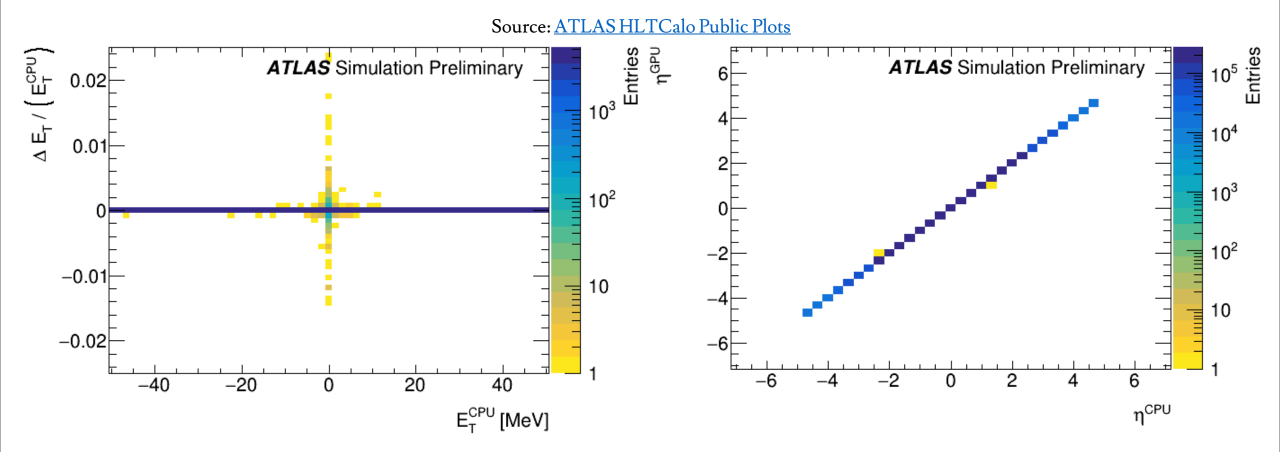
Topo-Automaton Clustering

- Growing, splitting and moments calculation fully implemented in the GPU using CUDA
- 100% agreement in cell assignment can be achieved between CPU and GPU with appropriate options (e. g. not using the two-Gaussian noise model)
 - Differences without these options are also fully understood (due to indeterminacies in the CPU and floating point accuracy issues, mostly)
- Basic cluster properties (e. g.: energy, η, φ) yield similar values (within floating point accuracy)
- Some cluster moments have greater differences due to accumulated and compounded floating point errors (there are calculated values that depend on calculated values that depend on calculated values...)
- The data structures used in the CPU part of the code cannot be used directly in the GPU, so we need to convert between the two representations



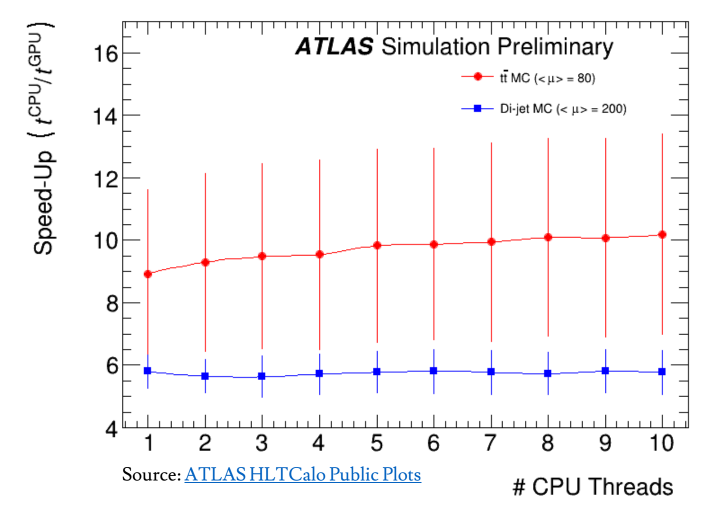
Comparison of the Cluster Properties

Relative error in the calculated transverse energy of the clusters as a function of the CPU reference value Comparison of the calculated pseudo-rapidity (η) of the clusters in the CPU and GPU implementations



Results show an **excellent agreement** between CPU and GPU, with only **floating point accuracy issues** caused by the **different order of operations** due to the inherently unpredictable timing of the parallel execution on a GPU.

Speed-up from GPU Acceleration in Relation to the CPU Implementation



We currently achieve a **speed-up of ~5.9 for di-jets, ~8.9 for** *tt*, considering all data conversions and transfers. The speed-up depends on the complexity of the event (number and size of the clusters), mostly due to CPU scaling.

Breakdown of GPU Execution Times

• Main bottleneck: converting the GPU data structures representing the clusters back to CPU-compatible structures

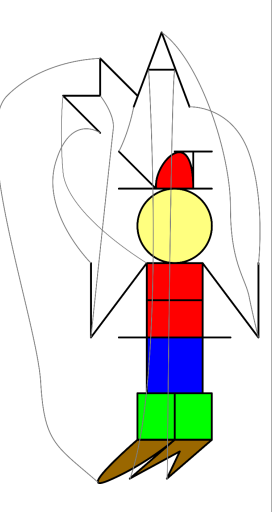
	C).		t - \bar{t} Events		Jet Events			
	Step	Time (µs)	s) Fraction of Total Time		Time (µs)	Fraction of Total Time		
Pr	e-Clustering Conversion	1441 ± 225	$9.24 \pm 1.58\%$		1128 ± 88	$13.24 \pm 1.14\%$		
P	re-Clustering Transfer	266 ± 8	8 $1.71 \pm 0.18\%$		248 ± 15	$2.92 \pm 0.30\%$		
	Cell Classification	61 ± 2	$15.76 \pm 1.77\%$		56 ± 3	$20.38 \pm 1.30\%$		
	Neighbour Pair Creation	159 ± 8	$40.68 \pm 3.23\%$	0.50 0.0504	114 ± 6	$41.45 \pm 1.98\%$	0.04 0.0507	
Growing	Tag Propagation	175 ± 46	$43.55 \pm 4.87\%$	$2.53 \pm 0.35\%$	106 ± 13	$38.17 \pm 2.59\%$	$3.24 \pm 0.25\%$	
	Total	396 ± 53	_		276 ± 16	_		
Post-G	rowing Property Calculation	74 ± 22	$0.47\pm$	0.14%	55 ± 2	$0.65 \pm 0.05\%$		
	Neighbour Pair Creation	409 ± 28	$29.23 \pm 2.73\%$		287 ± 14	$33.80 \pm 1.62\%$		
	Local Maxima Identification	88 ± 7	$6.25 \pm 0.53\%$		57 ± 3	$6.77 \pm 0.33\%$		
G. N.	Secondary Maxima Exclusion	229 ± 16	$16.48 \pm 2.25\%$	0.00 1.100	230 ± 14	$27.15 \pm 2.11\%$	0.05 0.5504	
Splitting	Main Tag Propagation	642 ± 194	$44.44 \pm 5.46\%$	$9.02 \pm 1.16\%$	236 ± 49	$27.53 \pm 3.54\%$	$9.97 \pm 0.57\%$	
	Finalization	50 ± 5	$3.60 \pm 0.31\%$		40 ± 3	$4.75 \pm 0.27\%$		
	Total	1417 ± 226	_		851 ± 67	_		
	Cluster Moments	1422 ± 134	$9.07 \pm 0.58\%$		889 ± 52	$10.43 \pm 0.51\%$		
Post-Clu	stering Transfer + Conversion	10679 ± 137	$67.77 \pm 2.59\%$		5094 ± 690	$59.27 \pm 2.26\%$		
	Total	15724 ± 1630		_	8565 ± 847	_		

- Potential improvements by **offloading ~3 ms of the conversion** to the GPU, but **the bottleneck would remain** given the constraints of **pre-existing CPU data structures** used in **other portions of the code**
- Ongoing effort to develop a **general solution** for an **Event Data Model** (EDM) with **CPU and GPU compatibility** requiring **minimal boilerplate**, with the possibility of providing an **API that matches pre-existing code**

EDM Developments

EDM Developments: Marionette

- <u>Marionette</u>: Memory Abstracted Representation with Interfaces in Objects Necessitating Extensively Templated Types EDM
- Goal: provide a more general solution to handle data structures that are meant to be usable on a CPU and a GPU (or a hardware accelerator in general) with a single source of truth
 - User-friendly, array-of-structs interface over the underlying struct-of-arrays
 - Arbitrarily extensible interface to enable compatibility with existing code
 - All the interface composition and data description handled at compilation time: no runtime polymorphism, the basic data structures are (almost) trivially copyable
 - No need to explicitly define CPU and GPU data structures and transfers "by hand", but specializations by the end user are fully possible
- Basic idea: decouple the description of the data to be stored from the way it which it will be laid out in memory: the data description is used to define the final data representation
 - The user simply provides a list of "properties" and specifies the layout
 - Strategic usage of macros allows injecting the name of the property as a part of the interface generated from it, for intuitive getter/setter behaviour: clusters[i].eta()
- Current status: the implementation works as intended and generates the same assembly as the equivalent handwritten structures, even for a complex example with GPU usage
 - Work in progress to reduce compilation times and further extend functionality



Marionette GPU Usage Example

```
//Declare two float properties with up to 256 entries, called 'energy' and 'time', associated with each entry of the collection.
MARIONETTE DECLARE PER ITEM PROPERTY SIZED(energy, Energy, 256, float);
                                                                              //provides x.energy() and x.setEnergy()
MARIONETTE DECLARE PER ITEM PROPERTY SIZED(time, Time, 256, float);
                                                                             //provides x.time() and x.setTime()
struct Foo : Marionette::InterfaceDescription::NoObject {
  template <class Final, class Layout> struct ObjectFunctions {
                                                                              //Define a property that adds a function
   int foo() const { return 42; }
                                                                              //to objects, but not the collection.
  };
};
struct Bar : Marionette::InterfaceDescription::NoObject {
  template <class Final, class Layout> struct ObjectFunctions {
                                                                       //These functions can refer to other functions
   float bar() const { return static_cast<const Final *>(this)->energy(); } //of the final collection or object via CRTP...
 };
};
//Define the list of properties to be used (the ones we defined earlier)
using ExampleProperties = Marionette::InterfaceDescription::PropertyList<Energy, Time, Foo, Bar>;
//Define the actual classes that hold memory as a struct-of-arrays with a fixed maximum size
template <class Context>
using OurCollection = OurCollectionBase<Marionette::LayoutTypes::DynamicStructInContext<Context, int>, ExampleProperties>;
using CPUCollection = OurCollection<Marionette::MemoryContexts::CUDAHostPinned>; //Pinned CPU memory for faster transfers
using GPUCollection = OurCollection<Marionette::MemoryContexts::CUDAStandardGPU>; //Normally allocated GPU memory
//Finally, the implementation:
CPUCollection coll(42);
                                                                              //Instantiate a collection of 42 elements
std::vector<float> desired times(50, 10.f);
                                                                              //Instantiate a collection of 42 elements
coll.time() = desired times;
                                                                              //coll.time() behaves to a vector of times
GPUCollection gpu coll = coll;
                                                                              //Copy-construct a new collection, held on the GPU
some kernel<<<4, 64>>>(Marionette::Collections::pass by value(gpu coll));
                                                                              //Pass that collection to a GPU kernel
coll = gpu coll;
                                                                              //Copy-assign back to the CPU collection
```

Summary and Future Efforts

Summary and Future Efforts

- Topo-Automaton Clustering **fully implemented** and working, for **cluster growing**, **cluster splitting** and **cluster moments calculation**, with configurability on a par with the CPU implementation (essentially, **drop-in replacement**)
- A very significant speed-up was found (factor of ~5.9 for di-jet events, ~8.9 for denser tt events)
 - This despite a significant portion of the GPU event processing time (60~70%) is spent in data conversions
 - Efforts to improve this bottleneck are under way, but it is a complex issue due to the well-established CPU structures and the nature of the underlying EDM approach
- Marionette may provide a general solution to mitigate the data structure conversion overhead
 - Some technical hurdles still to be overcome, but the current iteration of the Marionette framework works
 - Integration with the current implementation of Topo-Automaton Clustering to follow
- Preliminary work started on assessing the feasibility of performing at least some cluster calibrations on the GPU
- Lessons learned and experience gained from this development have fed back into general hardware acceleration-related development within ATLAS and in particular the ATLAS Trigger
- A final decision on using this approach in the ATLAS Trigger depends on a general technical assessment of the feasibility and/or performance of GPU-accelerated algorithms, being scheduled for next year
 - The approach is also being considered for offline reconstruction on grid sites where GPUs are available

Thank you for your attention!

Backup Slides

Conditions of the Benchmarking

- Samples correspond to two kinds of **Monte-Carlo simulated** events:
 - $t\bar{t}$ events: 3000 events, $\mu = 80$
 - **di-jet events**: 10000 events, $\mu = 200$
- Results were obtained on a remote server provided by the Brookhaven National Laboratory: GPU is a Tesla P100, CPU is a Xeon E5-2695 v4
- Time measurements were based on a per-thread clock
 - For a single thread, "any clock" would work
 - The CPU GPU comparison is a bit lopsided, though...
 - For more threads, timing and speed-up are representative, but throughput is a best-case estimate
 - o Essentially, we are assuming everything is always running in parallel
 - This is due to several limitations when trying to benchmark within the ATLAS software

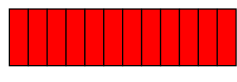
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	Total	1417 ± 226	_		851 ± 67	_		
	Cluster Moments	1422 ± 134	$9.07 \pm$	0.58%	889 ± 52	: 0.51%		
	Cluster Number Transfer	21 ± 1	$0.20 \pm 0.03\%$		17 ± 1	$0.33 \pm 0.04\%$		
	Cluster Info Transfer	250 ± 39	$2.35 \pm 0.27\%$		102 ± 20	$2.00 \pm 0.21\%$		
	Cluster Creation + Cell Info Transfer	395 ± 83	$3.68 \pm 0.46\%$		147 ± 24	$2.89 \pm 0.20\%$		
	Cell Cycle	2892 ± 444	$27.10 \pm 2.42\%$		1624 ± 117	$32.18 \pm 2.45\%$		
Importing	Cluster Ordering	192 ± 41	$1.79 \pm 0.28\%$	$67.77 \pm 2.59\%$	77 ± 15	$1.52 \pm 0.16\%$	$59.27 \pm 2.26\%$	
	Cluster Filling	2022 ± 336	$18.88 \pm 1.41\%$		944 ± 165	$18.46 \pm 1.18\%$		
	Moments Transfer	3.7 ± 0.5	$0.04 \pm 0.01\%$		3.8 ± 1.0	$0.07 \pm 0.02\%$		
	Moments Filling	4903 ± 697	$45.96 \pm 3.43\%$		2178 ± 389	$42.54 \pm 2.36\%$		
	Total	10679 ± 1377			5094 ± 690	_		
	Total	15724 ± 1630	-	_	8565 ± 847	_	_	

Topo-Automaton Cluster Growing – Anatomy of a Tag

High bit to distinguish valid tags from terminal and growing cells

Flag for preventing merges through seed cells (1 only in some edge cases with non-absolute value thresholds)



12 bit counter ($2^{12} - 1 - \#$ propagations)

Assumptions:

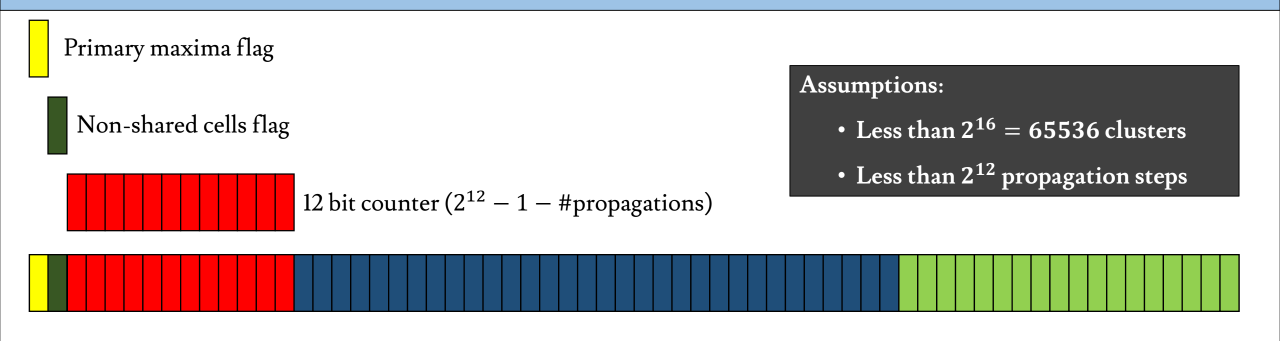
- Less than $2^{16} = 65536$ clusters
- Less than 2¹² propagation steps



Index of the seed cell

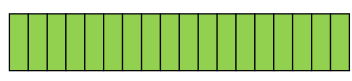


Topo-Automaton Cluster Splitting – Anatomy of a Tag





Index of the cell (or cluster index for original clusters)



Example Cell Indices

	00	01	02			03	0	05				06	07	08
_		09	0A	0B	ØC	ØD	04		0E	0F	10	11	12	
	13	14 15		1	6	17	18	19	1A	1	В	10	1 D	1E
	1F	20	15	2	1	22	23 24		25	26		1C	27	20
		29	2A	2B	2C	2D	2E	2F	31	32	33	34	35	28
-		36	37	38	39	3 A	3B 3C		3D	3E	3F	40	41	
		42				4	3	4	4			4	5	

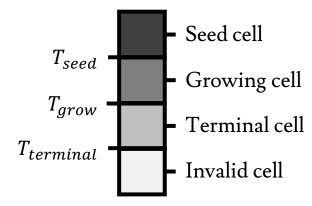
Example Cell Energies Bit Patterns

	3C	24	4B			64	<u>70</u>		05			6A	<u>7D</u>	63
		<u>6A</u>	63	6A	63	61			61	65	65	59	68	
	07	64			5	5D	59	59	5A	6	0		55	4D
	47	5B	5D	0	7	55	57	4D	4F	6	0	<u>6E</u>	64	C.F.
	47	<u>5C</u>	5B	4F	ØC	75	58	<u>5D</u>	5A	5C	5D	64	<u>67</u>	65
-		44	4C	2D	31	6A	5A	54	<u>5C</u>	57	<u>5F</u>	61	49	
		3F		0	0D		5A			7	С			

Topological Cluster Growing – Description

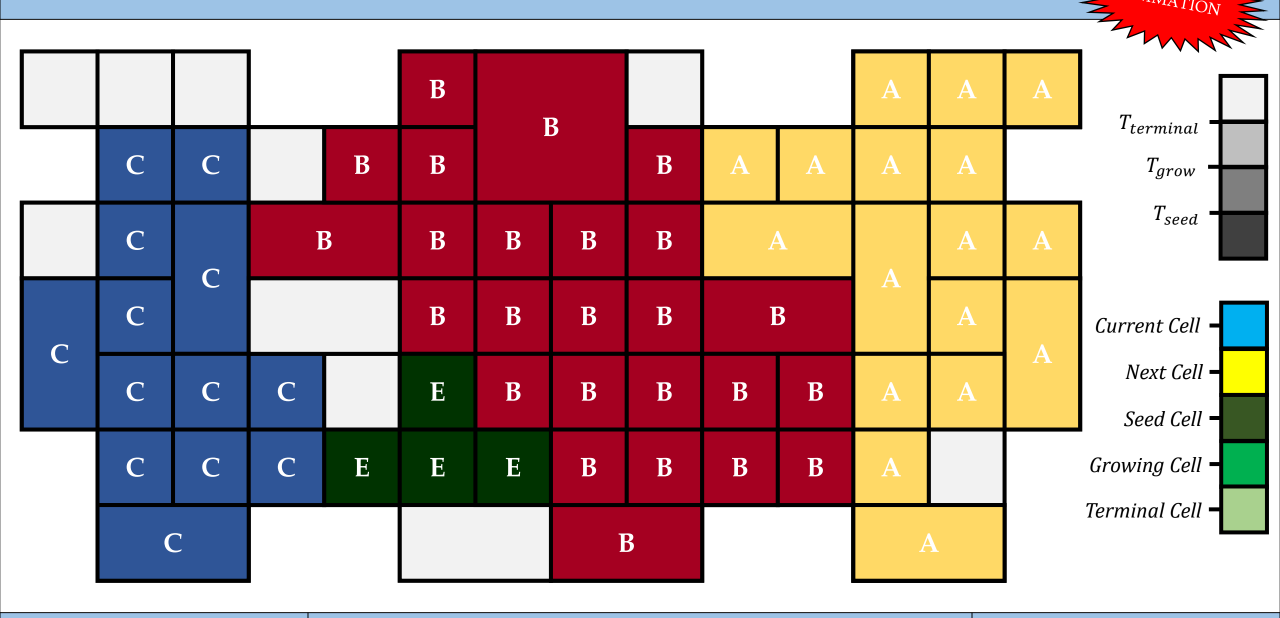
Cluster growing starts by **classifying the cells** using the **signal-to-noise ratio** of the energy deposition, according

to three thresholds:



- Seed cells are the **origin of the clusters** we construct, so we **sort them by the signal-to-noise ratio** and add them to the *list of current cells*, and we now **iterate until that list is empty**:
 - For every cell in the *list of current cells*, we will **check each of its direct neighbours**:
 - \circ If the neighbour does not belong to a cluster and its signal-to-noise ratio is above $T_{terminal}$,
 - ➤ Add it to the currently considered cell's cluster
 - \triangleright If its signal-to-noise ratio is above T_{grow} , add it to a *list of next cells*
 - \circ Else, if it already belongs to a cluster and its signal-to-noise ratio is above T_{grow} , merge the two clusters: all cells of the smallest cluster get added to the largest cluster
 - Once all cells have been checked, the *list of next cells* becomes the new *list of current cells*

Topological Cluster Growing – Example



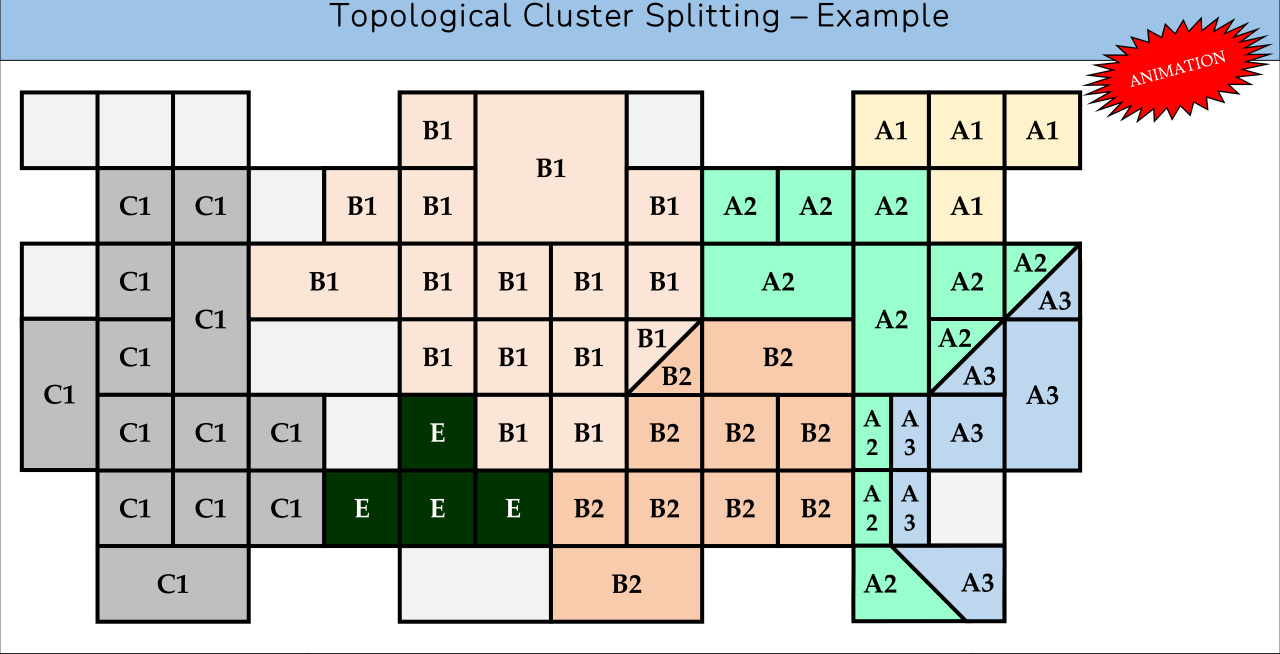
Topological Cluster Splitting – Description I

- Starting from the post-growing clusters, we **find the local maxima** of the deposited energy checking **only within the post-growing cluster** (and also taking into account some **thresholds in energy and number of neighbours**)
- Some calorimeter sampling layers give rise to **secondary local maxima**; secondary maxima that **overlap** with others through (next/prev)(InSamp/InCalo) neighbours are **excluded**
- Create a cluster for every local maximum, add the cells to the *list of current cells*, and **iterate until the list is empty**:
 - Sort the *list of current cells* by their energy
 - For every cell in the list, check each of its direct neighbours that is part of the same post-growing cluster:
 - If it does not belong to a cluster, add it to the current cell's cluster and to a list of next cells
 - Else, if it already belongs to a cluster, check if it is part of the *list of next cells*; if it is, **remove** it from that list, add it to a *list of next shared cells* and add it to the current cell's **cluster** as well
 - The *list of next cells* becomes the new *list of current cells*
 - Sort the *list of next shared cells* by **energy** and add it to the *list of shared cells*
- Add all elements of the *list of shared cells* to the (now empty) *list of current cells*

Topological Cluster Splitting – Description II

- Iterate again until the *list of current cells* is empty:
 - For every cell in the list, check each of its direct neighbours that is part of the same post-growing cluster:
 - o If it does not yet belong to a cluster, **add** it to both of the current cell's **clusters**, add it to the **list of shared** cells and add it to a **list of next cells**
 - The *list of next cells* becomes the new *list of current cells*
 - Sort the list of current cells by energy
- For every cell in the *list of shared cells*, **calculate the weight** of its contribution to each cluster: $w_1 = \frac{E_1}{E_1 + r E_2}$, $w_2 = 1 w_1$, with $r = e^{d_1 d_2}$ and d_i being the **distance** from the cell **to the centroid** of cluster i in units of typical shower scale (~ 5 cm)
- The resulting clusters are the **final clusters**
 - Post-growing clusters that had no local maxima are also part of the final clusters, unchanged by cluster splitting

Topological Cluster Splitting – Example



Cluster Moments Calculation

- While the calculation of cluster moments is **not the focus of the current discussion**, it is useful to provide a few remarks, especially to better **understand some of the comparisons** we will show later
- Many moments are weighted averages of cell properties (typically with $n \in \{1, 2\}$):

$$\langle \Xi^n \rangle = \frac{\sum_{c \in \text{cluster}: E_c > 0} w_c E_c (\Xi_c)^n}{\sum_{c \in \text{cluster}: E_c > 0} w_c E_c}$$

- Some moments depend on the **shower axis** \vec{S} (the **direction of flight** of the particle responsible for the shower):
 - Find the centroid of the cluster taking into account only cells with positive energy, $\vec{C} = (C_x, C_y, C_z)$
 - Define a matrix **M** such that its components are:

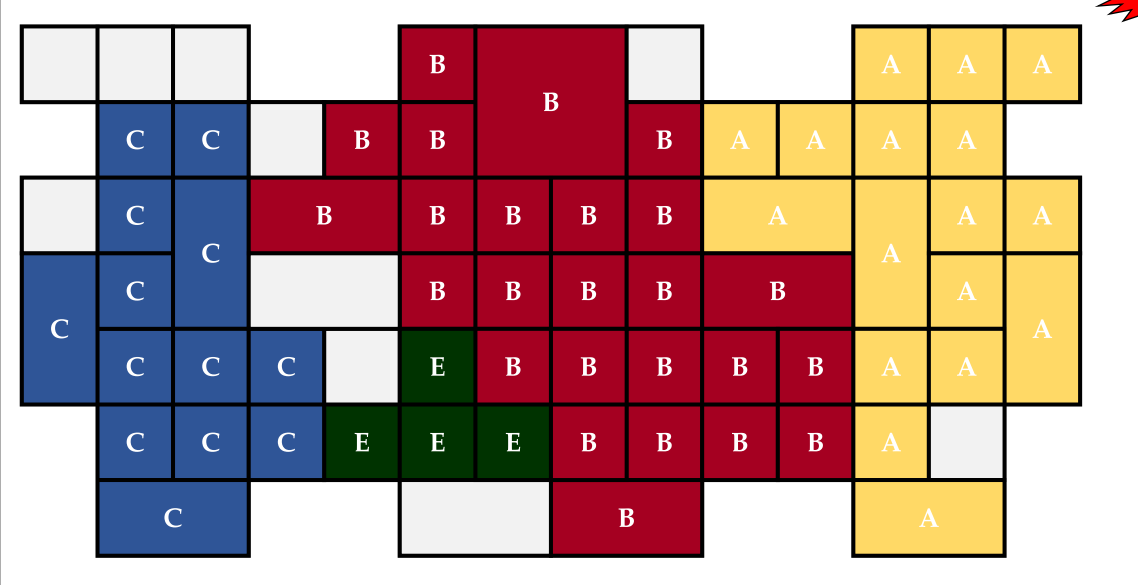
$$\mathbf{M}_{i,j} = \frac{\sum_{c \in \text{cluster}: E_c > 0} w_c^2 E_c^2 (x_i(c) - C_i) (x_j(c) - C_j)}{\sum_{c \in \text{cluster}: E_c > 0} w_c^2 E_c^2}$$

- The shower axis is the **normalized eigenvector** of **M** that has the **smallest angle** to the direction of \vec{C} , as long as that angle is smaller than a threshold (typically 20°) and the cluster has **at least 3 cells with positive energy**
- Otherwise, the shower axis is taken to be the direction of \vec{C} : $\vec{S} = \vec{C} / ||\vec{C}||$
- Some other moments (e. g. isolation, significance, second time) have different definitions

Topo-Automaton Cluster Growing – Description

- Classify each cell according to the signal-to-noise ratio, as in standard topological clustering:
 - Invalid, terminal and growing cells get assigned values that do not correspond to valid tags (0, 1 and $2^{63} 1$, respectively, though the choice is arbitrary as long as they are ordered and compare lower than any valid tags)
 - Seed cells are assigned a tag constructed as previously shown, with a counter for the cluster index being used (and incremented afterwards) to ensure the indices are sequential, and the corresponding entries of the seed cell index to cluster map and the cluster index to seed cell map are updated with the appropriate values
- Create two lists of pairs of neighbouring cells: the *list of growing pairs* for pairs of cells where both are growing or seed, and the *list of terminal pairs* where one is growing or seed and the other is terminal
 - Given that the **neighbourhood relations are not necessarily symmetric**, this is a list of **ordered pairs**: we will only consider **propagation in a particular direction** (by convention, from the second to the first cell of the pair)
- Keep iterating over all pairs of cells in the *list of growing pairs* until there are no tag changes:
 - If the second element of the pair is not part of a cluster, ignore this pair
 - Decrement the counter of the second element's tag and unset the cell merging flag to get the propagated tag
 - If the first element does not have a valid cluster tag, it gets the propagated tag (e. g. via an atomic maximum)
 - Else, if it is part of a different cluster, assign the highest of the two cluster indices to the entries of the seed cell index to cluster map and take the maximum between the entries of the cluster index to seed cell map
- For every pair of cells in the *list of terminal pairs*, propagate the tag from the second element to the first

Topo-Automaton Cluster Growing – Example



Topo-Automaton Cluster Splitting – Description I

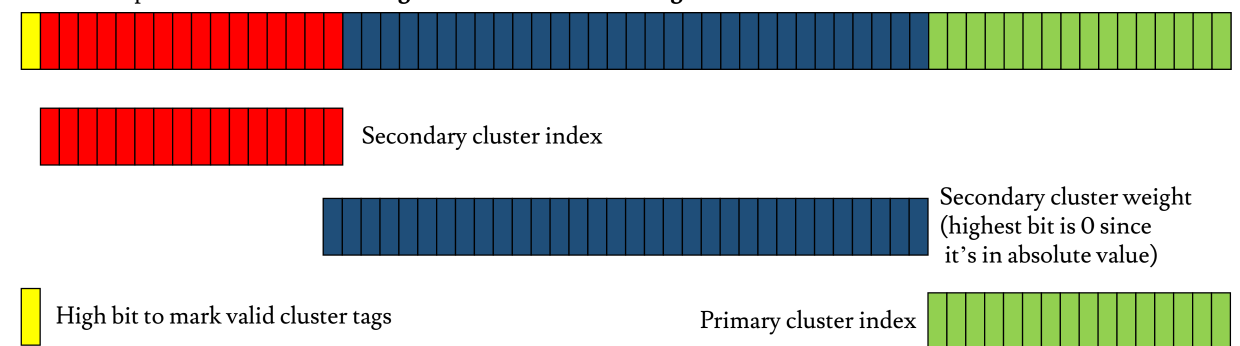
- Create four lists of ordered pairs of neighbours (similar to cluster growing) that will be used within the algorithm:
 - Pairs within the same post-growing cluster that will be used to expand the split clusters
 - Additional pairs used for checking the local maxima (due to certain neighbour options being ignored)
 - Pairs of next(InSamp/InCalo) neighbours regardless of the original post-growing cluster
 - Pairs of prev(InSamp/InCalo) neighbours regardless of the original post-growing cluster
- Use the first two lists of pairs to check for local maxima: exclude every cell that has a neighbour with equal or greater energy, check the remaining ones for the thresholds in energy and number of neighbours
- Local maxima are assigned a tag calculated as shown before (again using a counter and incrementing afterwards to get an index for every cluster) and the original cluster is stored in the cluster index to original cluster map; other cells belonging to clusters are assigned a tag that has the cluster index in the 18 lowest bits, and the first bit of the energy set if the post-growing cluster has local maxima, or all other bits and flags set if it does not
- Consider two separate sets of tags, one for next neighbours, the other for prev; every primary local maximum will be assigned the value of $2^{64} 1$, the secondary local maxima get their regular tags, all other cells are initialized to 0
- Keep iterating through the pairs of next and prev neighbours until there are no tag changes: if the second element of the pair has a non-zero tag, propagate it to the first
- Any **secondary maxima** that got their **tag replaced** in either set of tags should be **excluded** and their tags set to one that identifies a **cell belonging to the original cluster**, for all others **set the primary flag** (as there are **no further distinctions** between primary and secondary maxima in later stages of the algorithm)

Topo-Automaton Cluster Splitting – Description II

- Keep iterating until there are no tag changes:
 - Set a **reset counter** to 0
 - Iterate through the first list of pairs of neighbouring cells:
 - o If the second element of the pair is not part of a cluster created from a local maximum, ignore this pair
 - Obecrement the counter of the second element's tag (or set it to $2^{12} 1$ if the second element is a shared cell and the counter is higher than this value) and unset the primary flag to get the propagated tag
 - o If the first element does not have a valid cluster tag, it gets the propagated tag (e. g. via atomic maximum)
 - Else, if its reverse propagation counter is the same as the propagated tag's and it belongs to a different cluster, assign to it a tag that signals this is to become a shared cell: first flag and all bits of the counter set, second flag unset, and the energy and index the same as the propagated tag; also set the reset counter to the maximum between its current value and the reverse propagation counter of the original tag
 - Iterate through all cells that are part of a cluster created from a local maximum:
 - o If the cell's reverse propagation counter is lower than the reset counter, assign to it the tag of a cell that belongs to its original post-growing cluster (and thus not to a local maximum)
 - O Update the *cell index to cluster map* with the new cluster assignment (in the case of shared cells, the highest cluster index is in the most significant bits and the lowest in the least significant bits)
 - o Update the tag with the energy and index of this cell

Topo-Automaton Cluster Splitting – Description III

- Calculate the centroid of the split clusters: the absolute-energy-weighted sum of the non-shared cells' coordinates
- Assign appropriate weights to the contribution of the shared cells to each of the clusters:
 - As before, the weight is calculated by: $w_1 = \frac{E_1}{E_1 + r E_2}$, $w_2 = 1 w_1$, with $r = e^{d_1 d_2}$ and d_i being the distance from the cell to the centroid of cluster i in units of typical shower scale (~ 5 cm)
 - The choice of indices is such that $w_1 \le w_2$, to minimize potential precision loss
- We express the final cluster assignment with a different tag format:



Topo-Automaton Cluster Splitting – Example

