# Declarative Concurrent Programming with Join Calculus in Scala

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Lambda-Conf 2020 Global Edition

August 11, 2020

# Chemical Machine: a new hope

...and some new hype

#### In this talk:

- "join calculus" as a "Chemical Machine", without academic jargon
  - "λ-calculus" a small (but complete) programming language
  - ▶ "join calculus" a small (but complete) language for concurrency
    - $\star$  adds 2 features ("async mailbox" and "async function") to  $\lambda$ -calculus
    - ★ in the chemical metaphor: "molecule" and "reaction"
- Chymyst open-source implementation of Chemical Machine (Scala)
- examples of concurrent programs in Chymyst
  - ▶ implement anything in 10-15 lines of code
- o comparisons with Actor Model, "AWS Lambda", and Petri nets
- an extension for distributed programming: DCM

#### Not in this talk: academic theory

- Petri net theory,  $\pi$ -calculus, join calculus, joinads, formal semantics
- DCM defined within some formalism for distributed programming

# The chemical metaphor I. "Abstract" chemistry

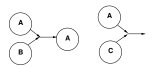
Real chemistry is asynchronous, concurrent, and parallel:

$$HCI + NaOH \rightarrow NaCI + H_2O$$

Want to run *computations* similarly to how chemical reactions run! Begin by formulating the execution model of "abstract chemistry":

- Abstract "molecules" float around in a "chemical reaction site"
- Certain kinds of molecules may combine to start a "reaction":

$$a + b \rightarrow a$$
  
 $a + c \rightarrow \emptyset$ 



- Program code defines molecules a, b, c, ... and chemical reactions
- At initial time, the code emits some molecules into the site
- The runtime system runs reactions concurrently and in parallel
  - ▶ A chemical simulation engine is easy to implement in any language

# The chemical metaphor II. Chemical Machine in a nutshell From abstract chemistry to computation

Translating the chemical metaphor into a model of computation:

- Each molecule carries a value ("concurrent data")
- Each reaction computes new values from its input values
- Some molecules with new values may be emitted back into the reaction site

```
site(
go { case a(x) + b(y) \Rightarrow
val z = f(x, y); a(z) \},
go { case a(x) + c(\_) \Rightarrow
println(x) }
```

When a reaction starts: input molecules disappear, new values are computed, output molecules are emitted

Reactions are functions from input values to output values

Need to learn how to "think in molecules and reactions"

# Programming the Chemical Machine using Chymyst

How I learned to forget semaphores and to love concurrency

Molecule emitters are values of type M[A]:

```
val count = m[Int]; val inc = m[Unit]
```

Molecules are emitted by calling the emitter's apply() method:

```
count(0); inc(); inc();
```

Reactions are values of type Reaction:

```
val r: Reaction = go { case count(x) + inc(_) \Rightarrow count(x + 1) }
```

- A reaction may be any partial function with a single case clause
- Emitters for all molecules (e.g., count, inc) must be already defined

Reactions are "activated" as a group, using a "reaction site":

```
site(r1, r2, r3)
```

- Molecules may be emitted after activating their consuming reactions
- A site *must* include all reactions that contend on an input molecule

Programs declare molecules and reaction sites, then emit initial molecules

The Chymyst runtime will run the simulation

## Example: throttling

Throttle emitting molecules s(x) with minimum allowed delay of delta ms

```
def throttle[X](s: M[X], delta: Long): M[X] = {
 val r = m[X]
 val enable = m[Unit] // Molecule is confined to the local scope.
 site(
  go { case r(x) + enable(_) \Rightarrow
        s(x)
        Thread.sleep(delta)
        enable()
 enable() // Enable emitting 's' initially.
 {f r} // Outside scope will be able to emit 'r'.
```

- No threads/semaphores/locks, no mutable state
- External code may emit r(x) at will, and s(x) is then throttled
- External code cannot emit enable() because of local scope

Implementations in Akka, in Monix, and ZIO: > 50 LOC each

## Example: throttling test

Define a molecule p(...) that we will emit too quickly, and a reaction just to see when that molecule has been emitted

```
val p = m[Int]
site(
   go { case p(x) ⇒ println(s"Got $x at time ${LocalDateTime.now}") }
)
val t = throttle(p, 1000L)
t(10); t(20); t(30);
scala> Got 10 at time 2020-08-10T15:33:05.921
Got 20 at time 2020-08-10T15:33:06.914
Got 30 at time 2020-08-10T15:33:07.920
```

## Example: cancellation

Cancel emitting molecules x(...) after a molecule cancel() is emitted

```
def cancellation[A](x: M[A]): (M[A], M[Unit]) = {
  val cx = m[A] // Cancellable version of 'x'.
  val cancel = m[Unit]; val enable = m[Unit]
  site(
    go { case cx(a) + enable(_) ⇒ x(a); enable() },
    go { case enable(_) + cancel(_) ⇒ },
  )
  enable() // Enable emitting 'x' initially.
  (cx, cancel) // Outside scope will be able to emit these.
}
```

- Emitting cancel() many times has no further effect
- External code cannot emit enable() because of local scope

Users cannot implement this in Monix or ZIO (cancellation is provided)

# Example: async racing

For two molecules p(...) and q(...), determine which one is emitted first and report its value on a given molecule r(...)

```
def racing[A, B](r: M[Either[A, B]]): (M[A], M[B]) = {
  val p = m[A]; val q = m[B]
  val enable = m[Unit]
  site(
    go { case p(a) + enable(_) ⇒ r(Left(a)) },
    go { case q(b) + enable(_) ⇒ r(Right(b)) },
  )
  enable() // One copy of 'enable' initially.
  (p, q) // Outside scope will be able to emit these.
}
```

- External code will emit p(...) and q(...), will get r(...) back
- External code cannot emit enable() because of local scope

Users cannot implement this in Monix or ZIO (racing facility is provided)

# Additional feature: blocking molecules

A blocking molecule has type B[A, R] and receives a reply value of type R:

```
val fetch = b[Unit, Int] // Blocking molecule's emitter.
val result = m[Int] // A non-blocking molecule emitter.
site(
  go { case fetch(_, reply) + res(b) ⇒ reply(b) }
) // 'reply' is a ReplyEmitter[Int] defined within reaction scope.
result(123)
val x = fetch() // Blocking call, will set x = 123.
```

Blocking emitters are a convenience, do not increase expressive power

### Example: racing with blocking molecules

Given two blocking molecules p(x:X) and q(y:Y), create a new blocking molecule r(...), to determine which one returns first and return the corresponding value of type Either[X, Y]

```
def racing[X,Y](p: B[Unit,X], q: B[Unit,Y]): B[Unit, Either[X,Y]] = {
   val get = b[Unit, Either[X, Y]]
   val res = b[Unit, Either[X, Y]]
   val left = m[Unit]; val right = m[Unit]
   val done = m[Either[X, Y]]
   site(
     go { case res(_, reply)
⇒ left(); right(); reply(get()) },
     go { case right(_{-}) \Rightarrow done(Right(_{+}())) },
     go { case get(\_, r) + done(x) \Rightarrow r(x) },
 res // Outside scope will be able to emit this.
```

• External code will call r(...), which calls p(...) and q(...)Users cannot implement this in Monix or ZIO (racing facility is provided)

## Example: map/reduce

#### A simple map/reduce implementation:

```
val c = m[A] // Initial values have type 'A'.
val d = m[(Int, B)] // B' is a commutative monoid.
val res = m[B] // Final result of type 'B'.
val fetch = b[Unit, B] // Blocking emitter.
site(
 // "map"
  go { case c(x) \Rightarrow d((1, long\_computation(x))) },
 // "reduce"
  go { case d((n1, b1)) + d((n2, b2)) \Rightarrow
  val (newN, newB) = (n1 + n2, b1 | + | b2)
   if (newN == total) res(newB) else d((newN, newB))
  },
  go { case fetch(\_, reply) + res(b) \Rightarrow reply(b) },
(1 to 100).foreach(x \Rightarrow c(x))
fetch() // Blocking call will return the final result.
```

Compare with the Akka implementation here (100+ LOC)

# Example: parallel merge-sort

#### Reactions can be recursive

Complete Chymyst code: MergeSortSpec.scala

```
val mergesort = m[(Array[T], M[Array[T]])]
site(
  go { case mergesort((arr, sortedResult)) ⇒
    if (arr.length <= 1) sortedResult(arr)</pre>
      else {
        val sorted1 = m[Array[T]]
        val sorted2 = m[Array[T]]
        site( // Define a lower-level reaction site.
          go { case sorted1(x) + sorted2(y) \Rightarrow
             sortedResult(arrayMerge(x,y))
        val (part1, part2) = arr.splitAt(arr.length/2)
        // Emit lower-level mergesort molecules:
        mergesort(part1, sorted1) + mergesort(part2, sorted2)
1)
```

Implementation in Akka: 30 LOC for the same functionality

# Dining philosophers I. Declarative vs. non-declarative code

The paradigmatic example of concurrency, parallelism and resource contention

Five philosophers sit at a round table, taking turns eating and thinking for random time intervals



Problem: simulate the process, avoiding deadlock and starvation Solutions in various programming languages: see Rosetta Code

• The Chemical Machine code is purely declarative

# Dining philosophers II. Implementation in Chymyst

Five Dining Philosophers implemented in 15 lines of code

```
Philosophers 1, 2, 3, 4, 5 and forks f12, f23, f34, f45, f51
     // ... definitions of emitters, think(), eat() omitted for brevity
     site (
        go { case t1(_) \Rightarrow think(1); h1() },
        go { case t2(\_) \Rightarrow think(2); h2() },
        go { case t3() \Rightarrow think(3); h3()  },
        go { case t4(\_) \Rightarrow think(4); h4() },
        go { case t5(\_) \Rightarrow think(5); h5() },
        go { case h1() + f12() + f51() \Rightarrow eat(1); t1() + f12() + f51() },
        go { case h2() + f23() + f12() \Rightarrow eat(2); t2() + f23() + f12() },
        go { case h3(_) + f34(_) + f23(_) \Rightarrow eat(3); t3() + f34() + f23() },
        go { case h4(_) + f45(_) + f34(_) \Rightarrow eat(4); t4() + f45() + f34() },
        go { case h5() + f51() + f45() \Rightarrow eat(5); t5() + f51() + f45() }
     t1() + t2() + t3() + t4() + t5()
     f12() + f23() + f34() + f45() + f51()
```

Source code: DiningPhilosophers.scala

For more examples, see the code repository (first-of, barriers, rendezvous, critical sections, readers/writers, Game of Life, elevators, etc.)

# Reasoning about code in the Chemical Machine paradigm

#### Reasoning about concurrent data:

- $\bullet$  Emit molecule with value  $\approx$  lift data into the "concurrent world"
- Define reaction ≈ lift a function into the "concurrent world"
- ullet Reaction site pprox container for concurrent functions and data
- ullet Reaction consumes molecules pprox function consumes input values
- ullet Reaction emits molecules pprox function returns result values

#### Reasoning about code:

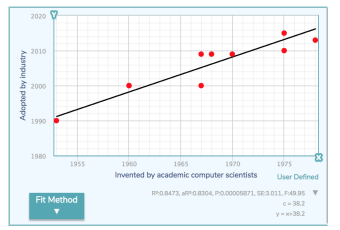
- What data do we need to handle concurrently? (Put it on molecules.)
- What computations consume this data? (Define as reactions.)

#### Guarantees:

- Molecule emitters and reactions are immutable values in local scopes
- Reaction sites are immutable once activated; can refactor to libraries
- Multiple input molecules are consumed atomically by reactions

# Chemical Machine paradigm to become mainstream in 2033

• The gap from academic invention to industry adoption is 38.2 years (infix math, continuations,  $\lambda$ -functions, OOP, CSP, map/reduce, Actor Model, constraint programming, DAG dataflow, Hindley-Milner types)



• The Chemical Machine paradigm was invented in 1995

# Current features of Chymyst

Experience with Chemical Machine programming is limited Some features promise to be useful:

- Blocking molecules with timeouts and timeout back-signalling
- Automatic pipelining of molecules (ordered mailboxes)
- Thread pools, thread priority control, graceful shutdown
- Facility to increase parallelism when using blocking code
- Errors in DSL are reported at compile-time or early run-time
  - Chymyst uses Scala macros but only to inspect code
  - Static analysis enables optimization and error reporting
- "Static" molecules with read-only access (similar to Akka "agents")
  - ► Forgetting to emit a molecule is #1 programmer error
- Logging, debugging, unit-testing facilities

#### Related frameworks: Petri nets

Workflow management: an approach based on Petri nets

- ING Baker a DSL for workflow management, based on Petri nets
- Process modeling and control ("elevator system" etc.)
- Business process management (BPM) systems

Chymyst implements a feature-rich version of Petri nets:

- Transitions admit arbitrary guard conditions and error recovery
- Transitions carry values, reactions are values, can be nested
- Nondeterministic, asynchronous, parallel execution

Any Petri net model is straightforwardly translated into a CM program

#### Chemical Machine vs. Amazon AWS Lambda

#### How AWS $\lambda$ works:

- wait for an event that signals arrival of input data
- run a computation whenever input data becomes available
- the computation is automatically parallelized, data-driven
- writing the output data will create a new event
- Modify the AWS $\lambda$  execution model by adding new requirements:
  - a Lambda should be able to wait for several *unrelated* events
  - several Lambdas may contend *atomically* on shared input events
- With these new requirements, AWS $\lambda$  becomes "AWS $\pi$ " a model of unrestricted concurrency
  - (Implementation on AWS could be tricky)

#### Chemical Machine vs. the Actor model. I

Modify the Actor execution model by adding new requirements:

- when messages arrive, actors are auto-created, maybe in parallel
- actors may wait atomically for messages in *several* different mailboxes It follows from these requirements that...
  - Auto-created actor instances are stateless and invisible to user
  - User code defines mailboxes and computations that consume messages
  - Repeated messages may be consumed in parallel
  - Messages are sent to mailboxes, not to specific actor instances:

```
// Akka // Chymyst val a: ActorRef = ... receive(x) \Rightarrow ... go { case a(x) \Rightarrow ... } val b: ActorRef = ... receive(y) \Rightarrow ... go { case b(y) + c(z) \Rightarrow ... } a ! 100 b ! 1; b ! 2; b ! 3 b ! 3; c("hello");
```

- All data resides on messages in mailboxes, is consumed automatically
- Mailboxes and computations are values, can be sent on messages

Any Actor program can be straightforwardly translated into CM

#### Chemical Machine vs. Actor model. II

- ullet reaction pprox function body for an (auto-started) actor
- ullet emitted molecule with value pprox message with value, in a mailbox
- molecule emitters ≈ mailbox references

#### Programming with actors:

- user code creates and manages explicit actor instances
- actors typically hold mutable state and/or mutate "behavior"
  - reasoning is about running processes and the data sent on messages

#### Programming with the Chemical Machine:

- processes auto-start when the needed input molecules are available
- many reactions may start at once, with automatic parallelism
  - user code does not manipulate references to processes
    - ★ no state, no supervision, no lifecycle, no "dead letters", no routers
  - reasoning is only about the data currently available on molecules
    - ★ no reasoning about running processes having internal state

Chymyst code is typically 2x - 3x shorter than equivalent Akka code

#### Distributed Chemical Machine

Run concurrent code on a cluster with no code changes

- Declare some molecules as "distributed", of type DM[T]
- No other new language constructions are necessary!
  - early prototype in progress, as extension of Chymyst

#### Distributed map/reduce in 15 LOC:

```
implicit val cluster = ClusterConfig(???)
val c = dm[Int] ; val d = dm[Int] // distributed
val res = m[(Int, List[Int])] // local
val fetch = b[Unit, List[Int]]
site(
  go { case c(x) \Rightarrow d(x * 2) }, // "map" on cluster,
// "reduce" on the driver node only.
  go { case res((n, list)) + d(x) \Rightarrow res((n-1, s::list)) },
// fetch results
  go { case fetch(_, reply) + res((0, list)) ⇒ reply(list) }
if (isDriver) { // 'true' only on the driver node.
  Seq(1, 2, 3).foreach(x \Rightarrow c(x))
  res((3, Nil)); fetch() // Returns the result.
```

Comparison: Akka implementation of distributed map/reduce (400+ LOC)

#### Distributed cache in 10 LOC

 Mutable Map[String, String] with operations: put, get, delete implicit val cluster = ClusterConfig(???)

```
implicit val cluster = ClusterConig(!!!)
val data = dm[mutable.Map[String, String]]
val put = dm[(String, String)]
val get = dm[(String, M[Option[String]]]
val delete = dm[String]
site(
   go { case data(dict) + put((k, v)) ⇒ data(dict.updated(k, v)) },
   go { case data(dict) + get((k, r)) ⇒ data(dict); r(dict.get(k)) },
   go { case data(dict) + delete(k) ⇒ dict.remove(k); data(dict) }
)
if (isDriver) data(mutable.Map[String, String]())
```

• Comparison: Distributed cache in 100 lines of Akka

# Distributed peer-to-peer chat in 15 LOC

- Register user names in chat room
- Fetch list of users
- Send and receive text messages

```
implicit val cluster = ClusterConfig(???)
val users = dm[List[DM[String]]] // List of users' message emitters.
val carrier = dm[DM[String]] // Carries this node's message emitter.
val fetch = b[Unit, List[DM[String]]]
site(go { case users(es) + carrier(e) ⇒ users(e :: es) }
, go { case users(es) + fetch(_{-}, r) \Rightarrow users(es); r(es) } )
val peerName = ??? // Read from config on node.
val sender = new DM[String](peerName) // Assign unique molecule name.
site(go { case sender(x) \Rightarrow println(s"Peer $peerName reads $x")})
carrier(sender)
if (isDriver) users(Nil)
// Fetch list of users and send a message to Sergei if present.
fetch()
  .find(_.name == "Sergei")
  .foreach(sender ⇒ sender("hello"))
```

Comparison: Distributed chat in > 100 lines of Akka

# Reasoning in the Distributed Chemical Machine

#### Distributed computing is made declarative

- Determine which data needs to be distributed and/or concurrent
- Determine which computations will need to consume that data
- Emit initial molecules and let the DCM run

#### Pure peer-to-peer architecture:

- Distributed molecules may be consumed by any DCM peer
- All DCM peers operate in the same way (no master/worker)
- All DCM peers need to define the same distributed reaction sites
  - Non-DCM code may differ between peers
  - Code or configuration could designate DCM peer as a "driver" or have different roles

# Chemical Machine: implementation details

- Each reaction site has a scheduler thread and a worker thread pool
- Each molecule is "bound" to a unique reaction site
- Each emitted molecule is stored in a multi-set at its reaction site
- Each emitted molecule triggers a search for possible reactions
  - ▶ Reaction search proceeds concurrently for different reaction sites
- Reactions are scheduled on the worker thread pool
  - ▶ The thread pool can be configured per-reaction or per-site
- Scala macros are used for static analysis and optimizations
  - Automatically pipelined molecules
  - Simplify and analyze Boolean conditions
- Error analysis is also performed at early run time
  - Reaction site with errors remain inactive

# Distributed Chemical Machine: implementation details

- Each distributed molecule (DM) is bound to a unique reaction site
- Emitted DM data goes into the ZK instance
- Each DCM peer listens to ZK messages and checks for its DMs
  - Once a DM is found, its data is downloaded and deserialized
- On a DCM peer, each DM is identified with a unique local RS
  - Downloaded molecules are emitted into the local RS to run reactions
  - ▶ All DCM peers must run identical reaction code for DMs
- Each DCM peer acquires a distributed lock on its DMs
  - ▶ Lock is released once reaction scheduling is complete
- If a node goes down or network fails, molecules will be unconsumed
  - Another DCM peer will pick up these molecules later

#### Conclusions and outlook

- Chemical Machine = declarative, purely functional concurrency
  - ▶ Enough power to replace threads, semaphors, atomic vars, etc.
  - Similar to Actor Model, but easier to use and "more purely functional"
  - Significantly shorter code, easier to reason about
- An open-source Scala implementation: Chymyst
  - Static DSL code analysis (with Scala macros)
  - Industry-strength features (thread priority control, pipelining, fault tolerance, unit testing and debugging APIs)
  - Extensive documentation: tutorial book
  - ▶ Distributed Chemical Machine work in progress
- Promising applications:
  - Workflow management, BPM
  - Asynchronous GUIs
  - Distributed peer-to-peer systems