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► **To cite this version:**

Pablo Inostroza, Tijs van Der Storm. The TTC 2014 Movie Database Case: Rascal Solution . Transformation Tool Contest, 2014, L'Aquila, Italy. pp.155 - 159. hal-01110851

HAL Id: hal-01110851

<https://inria.hal.science/hal-01110851>

Submitted on 29 Jan 2015

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The TTC 2014 Movie Database Case: Rascal Solution*

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Rascal is a meta-programming language for processing source code in the broad sense (models, documents, formats, languages, etc.). In this short note we discuss the implementation of the “TTC’14 Movie Database Case” in Rascal. In particular we will highlight the challenges and benefits of using a functional programming language for transforming graph-based models.

1 Introduction

Rascal is a meta-programming language for source code analysis and transformation [2, 3]. Concretely, it is targeted at analyzing and processing any kind of “source code in the broad sense”; this includes importing, analyzing, transforming, visualizing and generating, models, data files, program code, documentation, etc.

Rascal is a functional programming language in that all data is immutable (implemented using persistent data structures), and functional programming concepts are used throughout: algebraic data types, pattern matching, higher-order functions, comprehensions, etc.

Specifically for the domain of source code manipulation, Rascal features powerful primitives for parsing (context-free grammars), traversal (visit statement), relational analysis (transitive closure, image etc.), and code generation (string templates). The standard library includes programming language grammars (e.g., Java), IDE integration with Eclipse, numerous importers (e.g. XML, CSV, YAML, JSON etc.) and a rich visualization framework.

In the following sections we discuss the realization of the TTC’14 Movie Database case study [1] in Rascal. We conclude the paper with some observations and concluding remarks. All code examples can be found online at:

<https://github.com/cwi-swat/ttc2014-movie-database-case>

2 Description of the solution

Representing IMDB in Rascal

As Rascal is a functional programming language, where all data is immutable, the IMDB models have to be represented as trees instead of graphs. If there are cross references in the model, these have to be represented using (symbolic or opaque) identifiers which can be used to look up referenced elements. We use an algebraic data type to model IMDB models:

*This research was supported by the Netherlands Organisation for Scientific Research (NWO) Jacquard Grant “Next Generation Auditing: Data-Assurance as a service” (638.001.214).

```

data IMDB = imdb(map[Id, Movie] movies, map[Id, Person] persons,
                set[Group] groups, rel[Id movie, Id person] stars);
data Movie = movie(str title, real rating, int year);
data Person = actor(str name) | actress(str name);
data Group = couple(real avgRating, Id p1, Id p2, set[Id] movies);

```

An IMDB model is constructed using the `imdb` constructor. It contains the set of `movies`, `persons`, `groups` and a relation `stars` encoding which movie stars which persons. Both `movies` and `persons` are identified using the opaque `Id` type. To model this identification, the `movies` and `persons` field of an IMDB model are maps from such identifiers to the actual movie resp. person. `Movies` and `persons` are simple values containing the various fields that pertain to them. The `Group` type captures couples as required in Task 2. A `couple` references two persons and a set of movies using the opaque identifiers `Id`.

Task 1: Synthesizing Test Data

Synthesizing test data creates values of the type `IMDB` as declared in the previous section. The process starts with an empty model (`imdb((), (), {}, {}1)`), and then consecutively merges it with test models for a value in the range $1, \dots, n$. Each test model in turn consists of merging the negative and positive test model as discussed in the assignment. As an example, we list the creation of the positive test model:

```

IMDB createPositive(int i) = imdb(movies, people, {}, stars)
  when movies := ( j: movie("m<j>", toReal(j), 2013) | j <- [10*i..10*i+5] ),
    people := ( 10*i: actor("a<10*i>"), 10*i+1: actor("a<10*i+1>"),
              10*i+2: actor("a<10*i+2>"), 10*i+3: actress("a<10*i+3>"),
              10*i+4: actress("a<10*i+4>") ),
    stars := {<10*i, 10*i>, <10*i, 10*i+1>, <10*i, 10*i+2>, <10*i, 10*i+3>,
             <10*i+1, 10*i>, <10*i+1, 10*i+1>, <10*i+1, 10*i+2>, <10*i+1, 10*i+3>,
             <10*i+2, 10*i+1>, <10*i+2, 10*i+2>, <10*i+2, 10*i+3>,
             <10*i+3, 10*i+1>, <10*i+3, 10*i+2>, <10*i+3, 10*i+3>, <10*i+3, 10*i+4>,
             <10*i+4, 10*i+1>, <10*i+4, 10*i+2>, <10*i+4, 10*i+3>, <10*i+4, 10*i+4>};

```

The function uses map comprehensions to create the `movies` and `people` fields, and a binary relation literal to create the `stars` relation. It then simply returns a value containing all those fields.

Task 2: Adding Couples

Task 2 consists of enriching IMDB models with couples: pairs of persons that performed in the same movie, once or more often. This transformation is expressed by updating the `couples` field with the result of the function `makeCouples`:

```

public map[int, set[int]] computeCostars(rel[Id movie, Id person] stars, int n){
  map[int star, set[int] movies] moviesPerStar = toMap(invert(stars));
  map[int movie, set[int] stars] personsPerMovie = toMap(stars);
  return toMap({<m, p> | <m, p> <- stars, size(moviesPerStar[p])>=3, size(personsPerMovie[m])>=n});
}
set[Group] makeCouples(model:imdb(movies, persons, groups,stars)){
  map[int movie, set[int] stars] costars = computeCostars(stars, n);

```

¹The syntax `()` indicates an empty map, whereas `{1:"a", 2:"b"}` represents a map with keys 1,2 and values "a","b".

```

map[tuple[int star1, int star2] couple, set[int] movies] couples = ();
for (int movie <- costars, int s1 <- costars[movie], int s2 <- costars[movie], s1 < s2) {
  couples[<s1, s2>]?{} += {movie}; }
return { couple(0.0, x, y, ms) | <x, y> <- couples, ms := couples[<x, y>], size(ms) >=3 };
}

```

The `makeCouples` function first converts the binary relation `stars` to a map (`costars`) from movie Id to set of person Ids, filtering out some irrelevant elements by calling the `computeCostars` function. The central `for` loop iterates over all movies and all combinations of two actors and adds the movie to a table maintaining the set of movies for all couples (a map taking tuples of person Ids to sets of movie Ids). The side condition `s1 < s2` ensures we don't visit duplicate or self combinations. The question mark notation initializes a map entry with a default value, if the entry did not yet exist. In the final statement, a set of Groups is returned containing all couples which performed in 3 or more movies.

Task 3: Computing Average Ratings for Couples

As can be seen in the previous section, the average rating field of couples is initialized to 0.0. In this task we again transform an IMDB model, this time enriching each couple with its average rating of the movies the couple co-starred in. The following function performs this transformation:

```

IMDB addGroupRatings(IMDB m) = m[groups=gs]
  when gs := { g[avgRating = mean([ m.movies[x].rating | x <- g.movies ])] | g <- m.groups };

```

The `groups` field of the model `m` is updated with a new set of groups, as created in the `when`-clause of the function. The new set of groups is created using a comprehension, updating the `avgRating` field of each group. The average is computed based on the list of ratings obtained from the movies contained in `m` that are referenced in the group `g`.

Extension Task 1: Top 15 Rankings

The object of the extension Task 1 is to compute top 15 rankings based on the average ratings or number of movies a couple participated in. To represent rankings, we first introduce the following type alias:

```

alias Ranking = lrel[set[Person] persons, real avgRating, int numOfMovies];

```

A ranking is an ordered relation (`lrel`) containing the co-stars, the average rating and the number of movies of a couple. To generate rankings in this type, we create a generic function that takes the number of entries (e.g., 15), an IMDB model, and a predicate function to determine ordering of groups. This last argument allows to abstract over what the ranking is based (e.g., average rating or number of movies).

```

Ranking rank(int n, IMDB m, bool(Group, Group) gt) =
  take(n, [<{m.persons[x] | x <- getPersons(g)}, g.avgRating, size(g.movies)>
    | Group g <- sort(m.groups, gt)]);

```

Again, this function employs a comprehension to create a ranking, iterating over the groups in the IMDB model, sorted according to the predicate `gt`. For each person in the group (extracted using `getPersons`), the actual person value is looked up in the model (`m.person[x]`). The `take`, `size` and `sort` functions are in the standard library of Rascal.

The actual top 15 rankings are then obtained as follows:

```
Ranking top15avgRating(IMDB m) = rank(15, m, greaterThan(getRating));
Ranking top15commonMovies(IMDB m) = rank(15, m, greaterThan(getNumOfMovies));
```

The last argument to `rank` is constructed using a higher-order function, `greaterThan`, which constructs comparison functions on groups based on the argument getter function (i.e. `getRating` and `getNumOfMovies`). So in the first case, `rank` is called with a comparison predicate based on average ratings of groups, whereas in the second case, groups are ordered based on the number of shared movies in a group.

Extension Task 2: Generalizing groups to cliques

The extension task 2 consists of generalizing couples to arbitrarily sized cliques of people who co-starred in the same set of movies. A couple is a special case where the clique size is 2. To represent arbitrary cliques in the model, we modularly extended the `Group` data type (see Section 1.1) as follows:

```
data Group = clique(real avgRating, set[Id] persons, set[Id] movies);
```

This declaration states that the `clique` constructor is now a valid group value, in addition to `couple`.

Enriching an IMDB model with cliques follows the same pattern as enriching a model with couples. In fact the following function follows almost exactly the same structure as the function `makeCouples` described earlier:

```
set[Group] makeCliques(model:imdb(movies, persons, groups,stars), int n) {
  map[int movie, set[int] stars] costars = computeCostars(stars, n);
  map[set[int] clique, set[int] movies] cliques = ();
  for (Id movie <- costars, set[Id] s <- combinations(costars[movie], n)) {
    cliques[s]?{} += {movie}; }
  return {clique(0.0, s, ms) | s <- cliques, ms := cliques[s], size(ms) >= 3 };
}
```

Instead of iterating over pairs of actors explicitly, we now iterate over all combinations of size n using the helper function `combinations`, which generates all combinations of size n taking elements from the set `costars[movie]`.

Extension Task 3 & 4 These are the same as Task 3 and Extension Task 1, respectively, but intended for cliques instead of couples. It is not necessary to address these new cases in particular, because the code is polymorphic over groups. Only in the case of Extension Task 4, the `getPersons` accessor has to be extended to accommodate the new `clique` constructor defined in Extension Task 2:

```
set[Id] getPersons(clique(_, set[Id] ps, _)) = ps;
```

3 Observations and Concluding Remarks

Rascal can be seen as a model-transformation system, but it has to be acknowledged that its functional nature poses certain challenges when compared to traditional model-transformation platforms:

- Since Rascal is based on immutable data, models have to be represented as (containment) trees with explicit cross-references. As a result, all model elements need to have an *identifier* and some transformations have to explicitly look up model elements, given their identity.

- For the same reason, model transformations feature non-destructive rewriting. That is, it is not possible to perform in-place updates. This has benefits for reasoning (locality), but might affect performance. In other words, in order to benefit from equational reasoning, there will be a compromise in terms of performance. To improve this situation, active research is being conducted with the aim of optimizing the implementation of immutable data structures in Rascal [4].

Looking back at the effort implementing the TTC'14 tasks we can observe that Rascal posed no problems for solving the tasks. The solutions are small and declarative. The size of the implementation is around 130 SLOC, including some helper functions, but excluding loading the model from XML which is another 38 SLOC. Although Rascal allows side-effects in (local) variables, with the exception of `makeCouples` and `makeCliques` none of the function use side-effects of any kind.

Another observation is that the tasks mostly involved querying the models and aggregating new results to enrich the model. In such cases, comprehensions are valuable features to create sets, maps, or relations of model elements. Rascal's built-in features for traversal and powerful pattern matching (e.g., deep pattern matching), were not (even) needed to perform most of the tasks in an adequate way.

Related to the nature of the tasks, the fact that cross references had to be represented and managed explicitly posed no problem. In all cases the top-level IMDB model was always available to perform the necessary reference lookups in the `movies` and `persons` tables. It is, however, conceivable that in the case of more complex transformations (in which the referential structure of a model needs to be changed), more administration would be required in order to keep referential integrity intact.

In terms of performance, and in absence of suitable benchmarks to compare, we can at this point only report on the observed behavior of our solution. As an example, we observed that extracting cliques of size 3 takes more than 1 hour on a 3.3Mb IMDB file². As it was shown in the code for computing couples and cliques, we improved the performance by filtering out some model elements before the actual couple/cliQUE computation (e.g. removing actors who performed in less than 3 movies). By doing so we could process the same file in 3 minutes.

Finally, we would like to emphasize that Rascal's module system proved its value. Some tasks could be implemented as modular extensions of earlier tasks, combining extension of data types (Extension Task 2) and extension of functions (Extension Tasks 3 and 4). In that way, it was possible to define generic behavior for the `Group` ADT which initially considered just couples, but was later on modularly extended to consider cliques too. Few additions to the original code were necessary, as the functionality was defined in a generic way so that new variants could be naturally handled via polymorphism.

References

- [1] Tassilo Horn, Christian Krause & Matthias Tichy (2014): *The TTC 2014 Movie Database Case*. In: *7th Transformation Tool Contest (TTC 2014)*, EPTCS.
- [2] Paul Klint, Tijs van der Storm & Jurgen Vinju (2009): *Rascal: A domain-specific language for source code analysis and manipulation*. In: *SCAM*, pp. 168–177.
- [3] Paul Klint, Tijs van der Storm & Jurgen Vinju (2011): *EASY Meta-programming with Rascal*. In: *Generative and Transformational Techniques in Software Engineering III*, Lecture Notes in Computer Science, Springer.
- [4] Michael Steindorfer & Jurgen Vinju (2014): *Code Specialization for Memory Efficient Hash Tries (Short Paper)*. In: *Generative Programming and Component Engineering GPCE 2014, Proceedings*.

²We ran our experiments on a laptop Apple MacBook Pro with Intel i7 CPU running at 2.9 GHz and 8GB memory.