

CDM

Fast Minimization

KLAUS SUTNER

CARNEGIE MELLON UNIVERSITY

FALL 2023



1 Partition Refinement

2 Hopcroft's Algorithm

3 Valmari-Lehtinen

Minimization is a good example where efficient computation forces one to think more carefully than in math alone.

Suppose you have a DFA and want to construct the corresponding minimal one.

There is a trivial brute-force approach that establishes computability: Enumerate all smaller DFAs and check for equivalence.

Implementing the quotient idea brute force produces $O(n^4)$. With a little more effort one can get down to $O(n^2)$ and often better (Moore's algorithm).

Breaking through the quadratic barrier is a bit more complicated.

Mathematical Thinking: behavioral equivalence. Once the concept of behavior is clear, there is a straightforward algorithm for minimization. And, it's even polynomial time.

Algorithmic Thinking: refinement of equivalence relations. A better algorithm is obtained by thinking clearly about computing with equivalence relations (Moore). The reward is a clean, quadratic time algorithm (which is often much better than quadratic).

Smart Algo Thinking: baby-steps vs. giant-steps. Now things get tricky: all sub-quadratic algorithms require a much more careful argument and deeper algorithmic methods. A bit of creative insight is required to get down to log-linear. And doing things elegantly and efficiently is quite difficult.

We will switch back and forth between two natural representations of the same concept.

Equivalence Relations

A relation $\rho \subseteq A \times A$ that is reflexive, symmetric and transitive.

Partition

A collection B_1, B_2, \dots, B_k of pairwise disjoint, non-empty subsets of A such that $\bigcup B_i = A$ (the blocks of the partition).

As always, we need to worry about appropriate data structures and algorithms that operate on these data structures.

Recall our abstract scenario: we have an equivalence relation $\rho \subseteq Q \times Q$ and an endofunction $f : Q \rightarrow Q$.

We want to find the coarsest refinement $\hat{\rho}$ of ρ that is compatible with f :

$$p \hat{\rho} q \Rightarrow f(p) \hat{\rho} f(q)$$

This is accomplished by repeated application of a **refinement operator** R_f :

$$p \rho_f q \Leftrightarrow f(p) \rho f(q)$$

$$R_f(\rho) = \rho \sqcap \rho_f$$

In other words: $\hat{\rho}$ is the fixed point of ρ under R_f .

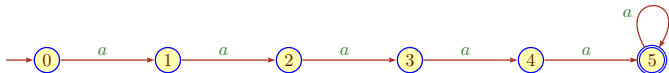
The refinement operator in Moore's algorithm works by representing all relations as canonical selector functions (aka int arrays) and scans the whole arrays for each step.

So each refinement step is $\Theta(n)$ (with good constants but still linear in n).

Quite often the algorithm uses fewer than n rounds, so the total time complexity may well be sub-quadratic.

Alas, there are cases when Moore requires $\Theta(n)$ rounds, producing quadratic running time overall.

Here is the standard example that demonstrates that Moore's algorithm may be quadratic: the minimal DFA for $\{a\}^{\geq k}$.



For this automaton, a single Moore round will split off only one state from the right end of block $D = \{0, 1, \dots, r\}$, at a cost of $\Theta(n)$ steps.

The split occurs only because of block $B = \{r + 1\}$, nothing else matters.

Critical Idea: Maybe we could get mileage out of trying to guide the refinement by single blocks, instead of blindly hitting the whole carrier set.

Suppose ρ is a partition of Q , and consider two blocks C and B . Let $f : Q \rightarrow Q$ be some endofunction.

We say that C **splits** B if

$$B \cap f^{-1}(C) \neq \emptyset \quad \text{and} \quad B - f^{-1}(C) \neq \emptyset.$$

In other words, $f(B)$ intersects both C and $Q - C$ and is not f -compatible yet: stopping the refinement process at this point would produce a nondeterministic machine. We need further refinement.

Let's define a new, more complicated refinement operator $\rho' = R_f(\rho, B, C)$ as follows:

$$p \rho' q \iff (p, q \notin B \wedge p \rho q) \vee \\ (p, q \in B \wedge (f(p) \in C \iff f(q) \in C))$$

In other words: outside of block B we keep the old ρ . Inside of B we check for C -equivalence of children.

So $R_f(\rho, B, C)$ is indeed a refinement of ρ : block B is split in two.

Proposition

- $R_f(\rho) \sqsubseteq R_f(\rho, B, C)$.
- $R_f(\rho) \neq \rho$ implies that $R_f(\rho, B, C) \neq \rho$ for some B and C .

In other words, we make no mistakes and we can't get stuck.

Proof.

$R_f(\rho)$ is $\prod_{C,B} R_f(\rho, B, C)$ and thus finer than each part.

If $R_f(\rho) \neq \rho$ there must be some block B and $p, q \in B$ such that $\neg(f(p) \rho f(q))$.

Let C be the block containing $f(p)$, done. □

Of course, from a complexity perspective this may not sound too promising: we are breaking one giant step into multiple baby steps. It is not unreasonable to suspect that this might even increase running time.

But: The baby steps provide much better control over the selection of the next refinement step: we can choose the blocks involved at will.

With a little effort this feature can be exploited sufficiently to speed up the whole process.

Since we are interested in the case where $f = \delta_a$ in a DFA, let us lighten notation a bit and write $a^{-1}C$ rather than $\delta_a^{-1}(C)$. Also, for $B, C \subseteq Q$, write

$$B/C$$

for the partition induced by splitting B via $a^{-1}C$: $B^+ = B \cap a^{-1}C$ and $B^- = B - a^{-1}C$.

Computing B/C should be very cheap.

1 Partition Refinement

2 **Hopcroft's Algorithm**

3 Valmari-Lehtinen

Suppose we have $a \in \Sigma$ and we want to insure compatibility with δ_a (see below for the general case). In addition, we have a 2-coloring χ of Q .

The algorithm maintains two data structures:

- a **partition** P of Q , representing the equivalence relation,
- a **split list** S with entries some of the blocks in the partition.

Both partitions are initialized by the coloring.

We refer to the blocks C in S as **active**: we will refine P by $a^{-1}C$.

The algorithm extracts an active block from the split list and tries to refine the blocks in the partition accordingly.

It then updates the split list in a clever way, and stops when the split list becomes empty.

It is convenient to initialize the partition and split lists according to some arbitrary coloring $\chi : Q \rightarrow \mathbf{2}$: this includes the usual final/non-final distinction but also accommodates other situations.

Initializing P is straightforward: every color produces one block.

The split list S only gets the smaller of the two blocks (we are dealing with the single-function case here, see below on how to handle larger alphabets).

initialize partitions P and S to the given coloring

while S not empty **do**

 extract C from S

 compute $\hat{C} = a^{-1}C$

foreach block B split by C **do**

$$B^+ = B \cap \hat{C}$$

$$B^- = B - B^+$$

 replace B by B^+ and B^- in P // update partition

if B is in S // update split list

then replace B by B^+, B^- in S

else replace B by the **smaller** of B^+, B^- in S

end

At first glance it may seem like we are not doing enough work: in the last case it feels like both B^+ and B^- , the parts of B obtained by splitting wrto the critical block C , should be added to the split list.

Otherwise we might miss out on splitting some other block D that is not split by B but split by B^+ or B^- . The algorithm might stop without having produced an f -compatible relation.

Fortunately, this cannot happen: in the critical case, if B^+ splits block D , then so does B^- , and the other way around. By picking the smaller of the two blocks, we get the desired speedup.

The following observation by Hopcroft makes this idea of “no-missing-work” more formal.

Let B be a block in a partition, split into B^+ and B^- . Let X be a subset of a block.

Proposition

$$X/B \cap X/B^+ = X/B \cap X/B^- = X/B^+ \cap X/B^-$$

Call a set $Z \subseteq Q$ of states **safe** for partition P (or simply P -safe) if $f^{-1}(Z)$ does not split any block in P .

We will show that the following assertion is a loop invariant:

$$\forall X \in P - S \exists A \subseteq S \left(X \cup \bigcup A \text{ } P\text{-safe} \right)$$

Clearly, this assertion holds before the loop ever executes when there are only two blocks in P (and one in S , the smaller of the two blocks).

As is customary, we indicate the value of a variable after one more execution of the loop-body by attaching a prime: so P' is the partition after one more round.

In the following we argue about the state of affairs at the end of a round.

So we need to show that

$$\forall X \in P - S \exists A \subseteq S \left(X \cup \bigcup A \text{ } P\text{-safe} \right)$$

implies that

$$\forall X \in P' - S' \exists A \subseteq S' \left(X \cup \bigcup A \text{ } P'\text{-safe} \right)$$

So assume we have some arbitrary block $X \in P' - S'$.

So X is an old block that has just been removed from the split list.

Since X is old, it has not been split in the last round, yet was removed from the split list.

The only way this can happen is that $C = X$ is the critical block. But then safety is a direct result of the construction.

This time we have an induction hypothesis: $Z = X \cup \bigcup A$ is P -safe for some $A \subseteq S$.

Case 2.1: $C \notin A$

In this case we can replace split blocks in A : replace X by X^+ and X^- (which are both active) and we have $Z = X \cup \bigcup A'$ is P -safe. It is easy to check that Z is also P' -safe.

Case 2.2: $C \in A$

Again, we can replace split blocks in A ; alas, we lose $C \notin S'$ to get $X \cup \bigcup A_0$. But no block $Z \in P'$ can be affected by this: it was already split wrto $f^{-1}(C)$ during the round.

So block X is new and was created by splitting a block Z in the last round.

Say, $X = Z^+$ and $Z \cup \bigcup A$ is P -safe for some $A \subseteq S$.

As before we can handle split blocks in A .

Case 3.1: $C \notin A$

Then $X \cup (\bigcup A' \cup Z^-)$ is P' -safe as in Case 2.1. Note that indeed $Z^- \in S'$ by construction.

Case 3.2: $C \in A$

In this case, $X \cup \bigcup A_0 \cup Z^-$ is P' -safe as in Case 2.2.

Each block in P is represented by a doubly-linked list.

We maintain an array of pointers to these lists for P and similarly for S . We also keep track of the cardinality of each block.

Furthermore, we have an array of pointers so that $\text{pos}[p]$ points to the list node containing p , plus information about the current block containing p .

The key part of each round is the computation of $\widehat{C} = a^{-1}C$. We may assume that $f^{-1}(p)$ has been precomputed for each state p . We can traverse C in time linear in $|C|$.

When a block B is hit for the first time, we start splitting it into two lists B^+ and B^- . If, in the end, $B^- = \emptyset$ we simply replace B by B^+ .

All this can be handled in time $O(|a^{-1}C|)$.

Let us say that a state p is **active** if $p \in \bigcup S$, **inactive** otherwise.

At level 0, at most half of all states are active.

Each state in the critical block C becomes inactive, but maybe reactivated later. Hence we can naturally assign activation levels 0, 1, 2, ... to all active states.

Recall that we only reactivate at most half the states in a block, so no state can be activated more than $\log n$ times. But then the total work computing pre-images of active states during the whole execution is just $O(n \log n)$.

Hence the running time of the whole algorithm is bounded by $O(n \log n)$.

This is a seminal paper that will bring tears to your eyes.

J. Hopcroft

[A \$N \log N\$ Algorithm for Minimizing States in a Finite Automaton](#)

STAN-CS-71-190

We modify the split list in the algorithm to contain entries

$$(a, C) \in S$$

where C is a block and $a \in \Sigma$: the intent is that we later refine via $a^{-1}C$.

Of course, (a, X) is **smaller** than (a, Y) if $|X| \leq |Y|$.

Note that we need to add (a, X) for all $a \in \Sigma$, which produces a running time of $O(kn \log n)$.

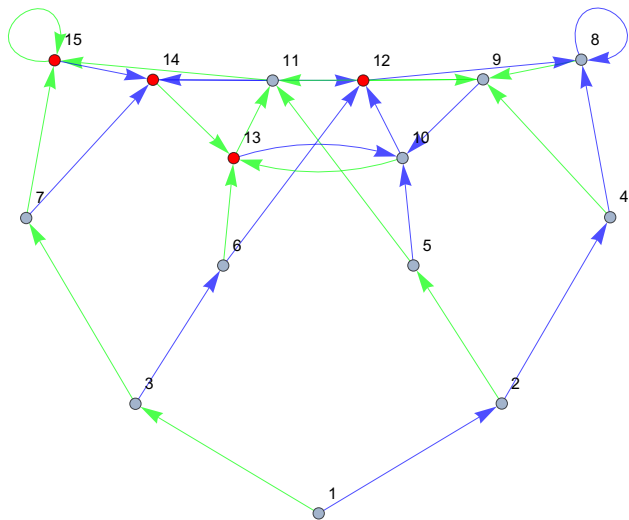
The following example uses a machine over alphabet $\{a, b\}$ with 15 states. The transition matrix is

	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
a	2	4	6	8	10	12	14	8	10	12	14	8	10	12	14
b	3	5	7	9	11	13	15	9	11	13	15	9	11	13	15

The final states are $\{12, 13, 14, 15\}$.

The following table shows the element extracted from the split list in the first column and the blocks in the second.

Note that split list entry (a, i) means: use the i th block in the current partition with respect to $f = \delta_a$.



$((12, 13, 14, 15), (1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11))$	$a, 1; b, 1$
$((14, 15), (1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11), (12, 13))$	$b, 1; a, 3; b, 3$
$((14, 15), (6, 7, 10, 11), (12, 13), (1, 2, 3, 4, 5, 8, 9))$	$b, 1; a, 3; b, 3; a, 2; b, 2$
$((15), (6, 7, 10, 11), (12, 13), (1, 2, 3, 4, 5, 8, 9), (14))$	$a, 3; b, 3; a, 2; b, 2; a, 5; b, 5$
$((15), (7, 11), (12, 13), (1, 2, 3, 4, 5, 8, 9), (14), (6, 10))$	$a, 3; b, 3; a, 2; b, 2; a, 5; b, 5; a, 6; b, 6$
$((15), (7, 11), (13), (1, 2, 3, 4, 5, 8, 9), (14), (6, 10), (12))$	$a, 5; b, 5; a, 6; b, 6; a, 7; b, 7$
$((15), (7, 11), (13), (3, 5, 9), (14), (6, 10), (12), (1, 2, 4, 8))$	$a, 5; b, 5; a, 6; b, 6; a, 7; b, 7; a, 4; b, 4$

In the first step, $a^{-1}(12, 13, 14, 15) = (6, 7, 10, 11, 14, 15)$ and we split both existing blocks.

This trace only shows steps where the partition changes. Note there are many “useless” steps at the end.

The algorithm is quite messy to implement correctly, as can be seen from the following papers:

D. Gries

[Describing an algorithm by Hopcroft](#)

Acta Informatica, 2 (1973) 97–109.

T. Knuutila

[Re-describing an algorithm by Hopcroft](#)

Theoretical Computer Science, 250 (2001) 333–363.

Exercise

Implement Hopcroft's algorithm, correctly.

Theorem (Hopcroft 1971)

Hopcroft's algorithm minimizes a DFA in $O(kn \log n)$ steps, where n is the state complexity of the DFA and k the size of the alphabet.

Unsurprisingly, the running time of Hopcroft's algorithm depends on the size k of the alphabet.

Knuutila pointed out in 2001, one can produce cubic (in n) running time when $k = n/2$ and a poor method of choosing the "smaller" block is used.

Note that Hopcroft's algorithm is nondeterministic in several ways.

- We can extract any element from the split list (e.g., could use a stack, queue, ...).
- Likewise we can place the new entries anywhere in the split list.
- When B^+ and B^- have the same size, we can pick either one.

None of these choices effect correctness, but they may well influence running time.

It should be noted that the algorithm often takes far fewer than $n \log n$ steps.

Given a reasonable implementation, every round is linear. It turns out to be quite difficult to construct inputs where the algorithm requires $\log n$ many rounds.

The best result known today is that there are some DFAs such that the algorithm takes $n \log n$ steps for a certain choice of active blocks in the main loop.

Alas, for these machines a different choice of active blocks results in linear running time.

- When is the running time $\Omega(n \log n)$ regardless of the chosen split list protocol? (A unary example is known where the execution sequence is essentially unique and reaches the log-lin bound.)
- What is the average complexity of Hopcroft's algorithm (average with respect to input automaton and/or split list protocol)?
- Is Hopcroft faster than Moore on average? For the uniform distribution, Moore has expected behavior $O(n \log n)$ and it may be that the constants are smaller (Bassino, David, Nicaud 2009).

1 Partition Refinement

2 Hopcroft's Algorithm

3 **Valmari-Lehtinen**

There are many examples where the number of transitions m in a partial DFA is much smaller than $k \cdot n$, k the size of the alphabet and n the number of states. This leads naturally to an algorithmic question:

Is there a $O(m \log n)$ minimization algorithm that deals directly with partial transition functions?

This would also nicely encapsulate problems with alphabet size in a parameter that really reflects the size of the data structure: unused symbols do not inflate the transition function.

The high-level logic is similar as in Hopcroft's algorithm: one maintains a partition of Q and tries to refine the partition by splitting with sets of the form

$$\delta_a^{-1}(B)$$

The choice of B is a bit more complicated, though.

Special care is taken to avoid unnecessary computation when preimages of blocks are computed. To this end, the algorithm also maintains and refines a second partition of the transitions.

As an implementation detail: the partition data structure is all array-based (unlike Hopcroft's algorithm).

Write $[n]_0$ for $\{0, 1, \dots, n - 1\}$.

Suppose we wish to maintain a partition of $[n]_0$. Keep track of r , the number of blocks, and maintain two maps (arrays)

$$\text{lo, hi: } [r]_0 \longrightarrow [n]_0$$

plus an array $P[n]$ such that

B_d , the block number d , is located in $P[\text{lo}[d], \text{hi}[d] - 1]$

We also have location and block-number maps

$$\text{loc: } [n]_0 \longrightarrow [n]_0$$

$$\text{bnum: } [n]_0 \longrightarrow [r]_0$$

such that $P[\text{loc}[p]] = p$, $\text{loc}[P[i]] = i$ and $p \in B_{\text{bnum}[p]}$.

It is convenient to subdivide the splitting process into three phases:

- Pre-splitting: initialize offset pointers $\text{mrk}[d] = \text{lo}[d]$ for all blocks, create empty hit list.
- Splitting: process a sequence of elements, swap each to the “marked” part $P[\text{lo}[d], \text{mrk}[d] - 1]$ of their respective blocks. If block B_d is encountered for the first time, add to hit list.
- Post-splitting: walk through blocks in hit list and update to maintain invariants.

It is straightforward to arrange the post-splitting phase so that whenever B splits into B_1 and B_2 , the larger part replaces B and the smaller part receives the higher block index (the current value of r).

As already mentioned, the minimization algorithm uses two PRDS:

- P : represents a partition of Q ; initialized to $(F, Q - F)$, as usual.
- T : represents a partition of the transitions; initialized to blocks containing all transitions with the same label.

The main loop of the algorithm looks like this:

```
foreach  $T$ -block  $C$  do  
  split blocks in  $P$  via  $C$   
  foreach  $P$ -block  $B$  do  
    split blocks in  $T$  via  $B$ 
```

New blocks are “appended” in both partitions, so the traversals end when all blocks have been processed.

```
pre-split  $P$   
foreach transition  $p \xrightarrow{a} q$  in block  $C$  do  
    mark source  $p$  in  $P$ -partition  
post-split  $P$ 
```

```
pre-split  $T$   
foreach state  $p$  in block  $B$  do  
    foreach transition  $t : q \xrightarrow{a} p$  do  
        mark  $t$  in  $T$ -partition  
post-split  $T$ 
```

Note that for the transition splitting operation one needs to be able to traverse all transitions with a fixed target.

To this end one precomputes two arrays

$$\text{trn} : [m]_0 \longrightarrow [m]_0$$

$$\text{fst} : [n] \longrightarrow [m]_0$$

such that for each state p

the transitions with target p are located in $\text{trn}[\text{fst}[p], \text{fst}[p + 1] - 1]$

This is easy via counting sort.

Recall that $\text{bnum}(p)$ is the block number of state p . By abuse of notation, $\text{bnum}(t)$ is the block number of transition t .

Proposition

- *States:* $\text{bnum}(p) \neq \text{bnum}(q)$ implies $\llbracket p \rrbracket \neq \llbracket q \rrbracket$
- *Transitions:* $\text{bnum}(s) \neq \text{bnum}(t)$ and $\text{lab}(s) = \text{lab}(t)$ implies $\llbracket \text{trg}(s) \rrbracket \neq \llbracket \text{trg}(t) \rrbracket$

Proposition

Let s, t be two transitions in the same T -block.

If $\text{src}(s)$ and $\text{src}(t)$ are in different P -blocks, then at least one of the blocks is unprocessed.

Proposition

Let p, q be two states in the same P -block and $s : p \xrightarrow{a} p'$, $t : q \xrightarrow{a} q'$ two transitions in different T -blocks. Then at least one of these blocks is unprocessed.

If $s : p \xrightarrow{a} p'$ and there is no a -transition with source q , then s is in an unprocessed T -block.

Theorem

The algorithm correctly minimizes a trim partial DFA in $O(m \log n)$ steps.

Proof.

It follows from the propositions that two states have the same behavior iff, upon completion, they are in the same P -block.

For running time, note that since we are dealing with a trim automaton, we have $n \leq m + 1$.

As in Hopcroft's algorithm, each state can be active at most $\log n$ times and, likewise, a transition can be active and at most $\log m$ times.

So the total running time is $O(m \log n)$.

This is another paper that will bring tears to your eyes, but for entirely different reasons.

Antti Valmari

[Fast brief practical DFA minimization](#)

Information Processing Letters 112 (2012) 213–217