Combinatorial approach to counting equivalence classes in 1D and 2D cellular automata

IZTOK JERAS^{1*}

Faculty of Computer and Information Science, University of Ljubljana, Slovenia

Received 17 December 2004; In final form 1 April 2005

There is much more theoretical research on 1D cellular automata compared to 2D. While not the only reason one of the arguments given is that the number of available rules for a common 2D neighborhood is too large for a comprehensive analysis. This issue can be mitigated by focusing on the smallest 2D neighborhoods (trid and quad) and by grouping rules into equivalence classes. This article gives an algebraic combinatorial solution to counting equivalence classes.

Key words: cellular automata, combinatorics, equivalence classes, congruence operations

1 INTRODUCTION

The focus of theoretical research is usually the simplest possible version of a system. For 1D CA elementary rules are the simplest which exhibit not trivial behavior. Rule equivalence has been used by many authors to organize 256 elementary CA rules into 88 equivalence classes. Representative rules from each class were then observed and described. This approach found the most interesting rules which were then further analyzed. Rule 110 is a good example.

The same approach was never applied to 2D CA. For example most studied 2D rule is Game of Life. It is based on a Moore neighborhood is binary cells.

^{*} email: iztok.jeras@rattus.info

There are $2^{2^9}=2^512$ possible rules, too many for a comprehensive analysis of the entire rule space. The von Neumann neighborhood with $2^{2^5}=2^32$ rules is still impractical, and there is no rule with this neighborhood on a binary lattice, which would receive as much attention as Game of Life. While this two neighborhoods were the focus of 2D CA research till recently, there are two simpler neighborhoods named quad and trid which were popularized by Tommaso Toffoli only in 2008 using the QUAD prize competition. The aim of the QUAD prize was to find a computation-universal 2-state cellular automaton on a 2-dimensional, 22-neighborhood. Edward Powley proved computational universality on an even smaller neighborhood, the trid.

The focus of my CA research are preimages. On 1D CA de Bruijn diagrams are the main tool used for this research. The size of this diagrams depends on the size of the rue space. While once the algorithms for calculating preimages are known, the ability to compute them is only limited by computer memory, the research of such algorithms usually requires the ability of the researcher to model such problems with mathematical notations like graphs and matrices. Only with 2D neighborhoods as small as quad and trid such representations become small enough for theoretical analysis with the help of visual representations.

The purpose of this article is to calculate the number of equivalence classes for a few small neighborhoods on a binary lattice (2D and 3D). The number of interesting rules can be further reduced by removing rules which can be represented with a smaller neighborhood. This are rules where one of the neighbors is not used in the local transition function, and higher order rules, those that can be represented with multiple steps of a smaller neighborhood rule.

??? Most research on CA was recently done by Hidenosuke Nishio [3], [2], [1], [4].

Tommaso Toffoli http://uncomp.uwe.ac.uk/automata2008/
files/quad.pdf

Edward Powley http://uncomp.uwe.ac.uk/automata2008/files/
quadprize_powley.pdf

2 FORMALIZATION

I will skip the full CA formalization and will focus on aspects observed in this article. What needs to be defined is the number of states of a cell, symmetries of the underlying lattice, the neighborhood shape, and the rule value.

Each cell c can be in one of k states, which are numbered starting with 0. There are k! permutations of the state set.

$$c \in 0, 1, ..., k - 1$$
$$|G^{per}| = k!$$

The underlaying lattice can have one or more dimensions. In 1D only reflection symmetry exists, while in 2D there is also a 3, 4 or 6 fold rotation symmetry. 3D or lattices with even more dimensions are not discussed here, although the same principles can be applied there.

Regardless of the underlying lattice only two reflections, unchanged and mirrored.

$$G^{ref} = (if, ref)$$
$$|G^{ref}| = 2$$

The number of rotations depends on the underlying lattice and is discussed for each neighborhood shape.

The neighborhood is defined by the number of cells m and its shape. Only the trid and quad 2D neighborhoods will be discussed. The neighborhood value is an ordered list of cell values. There are k^m possible neighborhood values.

$$n = (c_0, c_1, ..., c_{m-1})$$

The local transition function maps a neighborhood state into the next state of a cell.

$$f(n) = c_n$$

It is defined by as many cell values as there are neighborhood values.

$$r = (f(0), f(1), ..., f(k^m))$$

The relative position between a call and its neighborhood is not relevant for this article. But to understand the symmetries it is important to formalize the positions of cells inside a neighborhood.

2.1 1D

For 1D CA cells inside a neighborhood are numbered from left to right (see Figure 1).

2.2 2D trid

Due to symmetries all enumerations of neighborhood cells for trid are equivalent. Here enumeration starts in the left bottom corner and continues in a counter clockwise loop (see Figure 2).

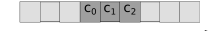


FIGURE 1

Elementary neighborhood on a linear lattice.

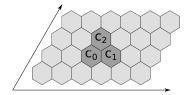


FIGURE 2

Trid neighborhood on a hexagonal lattice.

2.3 2D quad

To simplify the representation of rotations enumeration for quad also starts in the left bottom corner and continues in a counter clockwise loop (see Figure 3).

3 DEFINITION OF EQUIVALENCE

Two rules r_A, r_B are equivalent, if a transformation φ for the CA state exists, such that each CA current-next state pair (C^t, C^{t+1}) for the current rule gets transformed into a current-next state pair for the other rule.

$$r_A \sim r_B \Leftrightarrow \forall C_B = \varphi(C_A): F_B = \varphi(F_A): F_A(C_A^t) = C_A^{t+1}, F_B(C_B^t) = C_B^{t+1}: C_A^{t+1} = C_B^{t+1}$$

State transition graphs for equivalent rules are of the same shape, and a transformation exists which maps each state in one graph to a state at the same position in the graph for the other rule.

There are generalizations to this definition. For example a rule and it second order rule can be thought as equivalent. (rule performing the same global transition in one step as the original rule achieves in two steps.

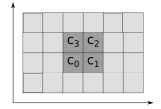


FIGURE 3 Quad neighborhood on a square lattice.

4 1D PROBLEM

Equivalence groups for elementary CA are long known. What is missing is a general algebraic solution.

4.1 Symmetries

There is a set of known symmetries or congruence operations on a given CA state. For 1D CA this are reflection and permutation.

Reflection:

$$G_{ref} = (id, ref)$$

 $|G_{ref}| = 2$

Permutation:

$$G_{per} = (id, per)$$
$$|G_{per}| = k!$$

4.2 Counting neighborhood values

The set of all distinct neighborhood values \mathbb{M} for a given neighborhood size n and number of cell states k is:

$$M = \{m :\}$$
$$|\mathbb{M}| = k^n$$

Neighborhood values can be organized into subsets depending on their invariance to congruence operations.

There are neighborhood values which are invariant to reflection. This are neighborhoods for which the left half is the mirror is the mirror image of the right half. There are as many such neighborhood values as there are possible values for a half sized neighborhood. For odd neighborhood sizes the middle cell is always a mirror image of itself.

$$\begin{split} \mathbb{M}_{\mathrm{ref}} &= \{ \forall n \in \mathbb{M} : \phi_{\mathrm{ref}}(n) = n \} \\ |\mathbb{M}_{\mathrm{ref}}| &= \left\{ \begin{array}{ll} k^{m/2+1} & \text{if } m \text{ is odd} \\ k^{m/2} & \text{if } m \text{ is even} \end{array} \right. \end{split}$$

There are no neighborhood values which are invariant to permutation.

$$\mathbb{M}_{per} = \{ \forall n \in \mathbb{M} : \phi_{per}(n) = n \} = \emptyset$$

$$|\mathbb{M}_{per}| = 0$$

There are neighborhood values which are invariant after both reflection and permutation have been applied.

$$\begin{split} \mathbb{M}_{\mathrm{ref,per}} &= \{ \forall n \in \mathbb{M} : \phi_{\mathrm{per}}(\phi_{\mathrm{ref}}(n)) = n \} \\ |\mathbb{M}_{\mathrm{ref,per}}| &= \left\{ \begin{array}{ll} 0 & \text{if } m \text{ is odd} \\ k^{m/2} & \text{if } m \text{ is even} \end{array} \right. \end{split}$$

4.3 Counting rules

Number of all rules |R|:

$$|R| = k^{|M|} = k^{k^n}$$

Rules which are invariant to reflection. The calculation is based on the number of reflection invariant neighborhood values. The local transition function can take any value for reflection invariant neighborhood values $M^{\rm ref}$ while for the rest $M\setminus M^{\rm ref}$ the same value must be used for the the neighborhood value and its reflection.

$$\begin{split} R^{\mathrm{ref}} &= \{\forall r \in R : \mathrm{ref}(r) = r\} \\ |R^{\mathrm{ref}}| &= k^{|M^{\mathrm{ref}}| + \frac{|M| - |M^{\mathrm{ref}}|}{2}} = k^{\frac{|M| + |M^{\mathrm{ref}}|}{2}} \\ |R^{\mathrm{ref}}| &= \left\{ \begin{array}{ll} k^{\frac{k^m + k^m/2 + 1}{2}} & \text{if } m \text{ is odd} \\ k^{\frac{k^m + k^m/2}{2}} & \text{if } m \text{ is even} \end{array} \right. \end{split}$$

Rules which are invariant to permutation.

$$R^{\mathrm{per}} = \{ \forall r \in R : \mathrm{per}(r) = r \}$$
$$|R^{\mathrm{per}}| = k^{|M|/k}$$

Rules which are invariant to reflection and permutation applied at the same time. First permutation limits free variables to $k^{|M|/k}$, then reflection limits them further by allowing a single free variable for all non reflective neighborhood values. TODO: check this for correctness.

$$R^{\text{ref\&per}} = \{ \forall r \in R : \text{per}(\text{ref}(r)) = r \}$$

k=2

$$|R^{\mathrm{ref\&per}}| = \left\{ egin{array}{ll} k^{1+1} & & \mathrm{if} \ m \ \mathrm{is} \ \mathrm{odd} \\ k^{k+1} & & \mathrm{if} \ m \ \mathrm{is} \ \mathrm{even} \end{array}
ight.$$

4.4 Counting equivalence classes

https://en.wikipedia.org/wiki/Burnside%27s_lemma Burnside's lemma is used to count equivalence classes:

$$|X/G| = \frac{1}{|G|} \sum_{g \in G} |X^g|$$

where G is

$$G_{ref} = (id, ref)$$

 $G_{per} = (id, per)$
 $G = G_{ref} \times G_{per} = (id, ref, per, ref \& per)$

$$|X/G| = \frac{1}{2 \times k!} (R + R^{ref} + (|G^{per}| - 1) * (R^{per} + R^{per\&ref}))$$

Result for elementary CA (k = 2, n = 3)

$$|X/G| = \frac{1}{4}(2^8 + 2^6 + 2^4 + 2^4) = 88$$

5 2D PROBLEM

5.1 Symmetries

In addition to reflection and permutation symmetries already present in 1D CA, 2D CA also exhibit rotation symmetry. The number of available rotations depends on the underlaying lattice.

3-fold: trid 4-fold: quad, Moore, von Neumann neighborhood 6-fold: hexagonal lattice

$$G_{rot} = (id, rot)$$

 $|G_{per}| =$

5.2 Trid

Counter clockwise rotations for 120°.

$$G_{rot} = (id = rot0, rot1, rot2)$$

 $|G_{rot}| = 3$

$$G = G_{ref} \times G_{per} \times G_{rot}$$
$$|G| = 12$$

$$|X/G| = \frac{1}{12} \left((2^8 + 2^6 + 2^4 + 2^4) + (2^4 + 2^2 + 2^6 + 2^4) + (2^2 + 2^2 + 2^6 + 2^4) \right) = 45$$

5.3 Quad

Counter clockwise rotations for 90°.

$$G_{rot} = (id = rot0, rot1, rot2, rot3)$$

$$|G_{rot}| = 4$$

$$G = G_{ref} \times G_{per} \times G_{rot}$$

$$|G| = 16$$

$$|X/G| = \frac{1}{16} \left((2^{16} + 2^8 + 2^{10} + 2^{10}) + (...) + (...) \right) = ...$$

6 CONCLUSION

With elementary cellular automata the approach was to observe the evolution of each rule and to further study the ones which exhibit interesting behavior. A similar approach can be taken with 2D trid and quad. With an appropriate filter this evolutions could be made into a video or a 3D image. An example filter would convert each 33 neighborhood pattern probabilities into a color palette. For trid a single person could do the review, while for quad crowd-sourcing could be used.

I hope will help further theoretical research into 2D CA.

		pe	per		ref		per,ref	
000	0	111	7	000	0	111	7	
001	1	110	6	100	4	011	3	
010	2	101	5	010	2	101	5	
011	3	100	4	110	6	001	1	
100	4	011	3	001	1	110	6	
101	5	010	2	101	5	010	2	
110	6	001	1	011	3	100	4	
111	7	900	θ	111	7	000	Θ	
rot1	rot1		rot1,per		rot1,ref		rot1,per,ref	
000	0	111	7	000	0	111	7	
010	2	101	5	001	1	110	6	
100	4	011	3	100	4	011	3	
110	6	001	1	101	5	010	2	
001	1	110	6	010	2	101	5	
011	3	100	4	011	3	100	4	
101	5	010	2	110	6	001	1	
111	7	000	θ	111	7	000	Θ	
rot2	rot2		rot2,per		rot2,ref		rot2,per,ref	
000	0	111	7	000	0	111	7	
100	4	011	3	010	2	101	5	
001	1	110	6	001	1	110	6	
101	5	010	2	011	3	100	4	
010	2	101	5	100	4	011	3	
110	6	001	1	110	6	001	1	
011	3	100	4	101	5	010	2	
111	7	000	0	111	7	000	θ	

TABLE 1 My caption

		per		ref		per,ref	
0000	0	1111	F	0000	0	1111	F
0001	1	1110	E	1000	8	0111	7
0010	2	1101	D	0100	4	1011	В
0011	3	1100	C	1100	C	0011	3
0100	4	1011	В	0010	2	1101	D
0101	5	1010	A	1010	A	0101	5
0110	6	1001	9	0110	6	1001	9
0111	7	1000	8	1110	E	0001	1
1000	8	0111	7	0001	1	1110	E
1001	9	0110	6	1001	9	0110	6
1010	A	0101	5	0101	5	1010	A
1011	В	0100	4	1101	D	0010	2
1100	C	0011	3	0011	3	1100	C
1101	D	0010	2	1011	В	0100	4
1110	E	0001	1	0111	7	1000	8
1111	F	0000	θ	1111	F	0000	θ

TABLE 2 My caption

REFERENCES

- [1] Hidenosuke Nishio. (2009). Automorphism classification of cellular automata. In Henning Bordihn, Rudolf Freund, Markus Holzer, Martin Kutrib, and Friedrich Otto, editors, Workshop on Non-Classical Models for Automata and Applications NCMA 2009, Wroclaw, Poland, August 31 September 1, 2009. Proceedings, volume 256 of books@ocg.at, pages 195–208. Austrian Computer Society.
- [2] Hidenosuke Nishio. (2010). Automorphissm classification of cellular automata. Fundam. Inform., 104(1-2):125–140.
- [3] Hidenosuke Nishio. (2012). A generalization of automorphism classification of cellular automata. *J. Cellular Automata*, 7(2):167–177.
- [4] Hidenosuke Nishio and Thomas Worsch. (2008). Changing the neighborhood of CA: local structure, equivalence and reversibility. In Andrew Adamatzky, Ramón Alonso-Sanz, Anna T. Lawniczak, Genaro Juárez Martínez, Kenichi Morita, and Thomas Worsch, editors, Automata 2008: Theory and Applications of Cellular Automata, Bristol, UK, June 12-14, 2008, pages 270–277. Luniver Press, Frome, UK.