Modeling and Controlling Parallel Tasks in Droplet-Based Microfluidic Systems

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Abstract—This paper presents general hardware-independent models and algorithms to automate the operation of droplet-based microfluidic systems. In these systems, discrete liquid volumes of typically less than 1 μ l are transported across a planar array by dielectrophoretic or electrowetting effects for biochemical analysis. Unlike in systems based on continuous flow through channels, valves, and pumps, the droplet paths can be reconfigured on demand and even in real time. Algorithms that generate efficient sequences of control signals for moving one or many droplets from start to goal positions, subject to constraints such as specific features and obstacles on the array surface or limitations in the control circuitry, are developed. In addition, an approach toward automatic mapping of a biochemical analysis task onto a DMFS is investigated. Achieving optimality in these algorithms can be prohibitive for large-scale configurations because of the high asymptotic complexity of coordinating multiple moving droplets. Instead, these algorithms achieve a compromise between high runtime efficiency and a more limited nonglobal optimality in the generated control sequences.

Index Terms—Digital microfluidic system, droplet-based microfluidic system, lab on a chip (LOC), parallel manipulation.

I. INTRODUCTION

DVANCES in microfabrication and microelectromechanical systems (MEMS) over the past decades have led to a rapidly expanding collection of techniques to build systems for the handling and analyzing of very small quantities of liquids (see, e.g., [1] and [2]). These microfluidic systems typically consist of submillimeter scale components such as channels, valves, pumps, and reservoirs, as well as application-specific sensors and actuators. Microfluidic devices hold great promise, for example, for novel fast, low-cost, portable, and disposable diagnostic tools. Applications include the massively parallel testing of new drugs, the on-site real-time detection of toxins and pathogens, and polymerase chain reaction (PCR) for deoxyribonucleic acid (DNA) sequence analysis. They usually operate with continuous flows of liquids, in analogy to traditional macroscale laboratory setups, and can integrate all

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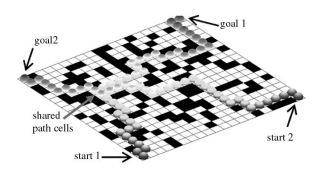


Fig. 1. Two droplets moving in parallel on a DMFS consisting of a 20×20 array with obstacle cells (marked black). The droplets start from cells (1,1) and (20,1) and move to cells (20,20) and (1,20), respectively. Change in droplet shading indicates the elapsed time. The droplets share cells (5,13) and (5,14) on their path but their coordinated schedule prevents any conflicts.

functionality into a complete lab on a chip (LOC) or biosystem on a chip (bioSOC).

More recently, an alternative LOC approach has gained momentum using individual droplets, with volumes usually in the submicroliter range. In these droplet-based microfluidic systems, droplets are generated, transported, merged, analyzed, and disposed on planar arrays of addressable cells; therefore, they are also sometimes called discrete or digital microfluidic systems, and conveniently abbreviated as DMFS. This architecture for microfluidic systems is attractive because of the following reasons: 1) greater flexibility—analyte handling may be reconfigured simply by reprogramming rather than by changing the physical layout of the microfluidic components; 2) high droplet speeds—reportedly up to 25 cm/s [3], [4]; 3) no dilution and cross-contamination due to diffusion and shear flow; and 4) the possibility for massively parallel operation.

The DMFS approach assumes that it is advantageous to shift complexity from microfluidic hardware to control software. Therefore, for a DMFS to live up to its promise, it must be accompanied by a complementary set of software tools such that its usage can be largely automated. This includes software that helps the user to map a biochemical analysis protocol onto a given DMFS; as a specific subproblem, algorithms that automatically plan and schedule routes for simultaneous droplet motion are required. Fig. 1 shows a schematic example where two droplets move in parallel across a DMFS while circumnavigating numerous obstacles. Developing the formalisms, models, and control strategies for such automated droplet manipulation tasks is the goal of this paper.

Processing large numbers of discrete droplets simultaneously on an integrated microchip indicates a similarity to electronic digital circuits, giving rise to microfluidic circuits [5], [6]. This analogy also suggests that algorithms for layout, routing, and scheduling of droplet paths in a DMFS are computationally expensive, i.e., NP-hard.

Thus, this paper is organized as follows. Section II reviews background material on DMFS hardware, and discusses related work in control algorithms. Section III introduces a formal DMFS model and problem specification. Section IV presents algorithms for coordinating parallel droplet motion on a DMFS and investigates tradeoffs between runtime efficiency and optimality. Section V extends these algorithms to allow for changes of the droplet type during DMFS operation and develops an approach to automatically transform a laboratory protocol into a sequence of DMFS tasks. Section VI concludes the paper with a summary and an outlook on future work.

II. RELATED WORK

Transferring a laboratory task such as DNA analysis, clinical diagnostics, or detection and manipulation of biomolecules into an LOC system is a complex endeavor that can involve multiple challenges: the design of microfluidic hardware including sensing and actuation mechanisms for liquid analytes; the use of specialized techniques and materials such as modification and functionalization of surfaces with monolayers or antibodies; and the development of algorithms for layout and control of massively parallel microfluidic circuits.

LOC design and manufacture has become an extensive research area with dedicated conferences (e.g., [7]) and journals (e.g., [8] and [9]). This paper, however, focuses on the software aspects, and assumes a device model that abstracts away from details of the physical implementation. Here, we discuss very briefly important aspects of droplet-based microfluidics that are relevant to motivate and justify our modeling assumptions.

A. Droplet Transport Techniques

The most successful conventional droplet-based system in the life sciences is arguably the fluorescence-activated cell sorter (FACS) [10]–[12], a machine that can sort droplets containing single cells at rates well above 100 kHz. It generates charged droplets, analyzes them in free flight via a laser fluorescence detection system, and sorts them accordingly via a modulated electrostatic field. LOC FACS systems exist but so far work at much lower processing rates [13]–[15].

In microscale LOC systems, droplets can be moved across a planar surface effectively with a variety of techniques, including electric fields (e.g., [3] and [16]–[19]), surface acoustic waves (e.g., [20]–[22]), thermocapillary and Marangoni effects (e.g., [23] and [24]), electrochemical surface modulation (e.g., [25]), conformational changes in molecular surface layers (e.g., [26]), or gradients in surface chemistry (e.g., [27] and [28]), and texture (e.g., [29] and [30]). For this paper, droplet transport with high speed, accuracy, and full software control is essential, making electric fields the most suitable approach; hence, we briefly discuss the two main techniques in this realm, i.e., dielectrophoresis (DEP) and electrowetting [31], [32].

1) Dielectrophoresis: In DEP, neutrally charged objects are first polarized by an electric field and then experience a net

force due to the field. This force can only be nonzero if a field gradient exists, i.e., the positively and negatively polarized regions of the object occupy areas of different field strengths. If the object has stronger polarization than the surrounding medium, then it is pulled toward the areas of higher field strength (this is called positive DEP), but if the surrounding medium has higher polarization, then the object is pushed toward areas of lower field strength (negative DEP). DEP can be considered the electrostatic analogy of induced magnetism. Common examples for DEP are charged clothes that attract (neutral) lint particles. More information on DEP can be found, e.g., in [33]. A DMFS system employing DEP with more than 10 000 array elements was demonstrated in [34].

2) Electrowetting: Electrowetting on dielectric (EWOD) exploits the decrease of contact angle that an aqueous droplet on a dielectric surface experiences when exposed to an electric field. If the field is localized at only one side of the droplet, then the difference in contact angle causes a pressure differential in the droplet, which drives it toward the region of higher field strength. Electrowetting and its applications in microfluidics have been investigated by several groups, including [3], [16], [17], [35], and [36].

B. Droplet Transport Planning and Scheduling

Finding the optimal plan to generate, store, move, merge, split, and dispose multiple droplets on a DMFS combines general path planning and scheduling with the more application-specific task of analyte droplet handling. Various researchers have studied parts of the overall problem and have shown important results on algorithmic solutions and their computational complexity.

One possible approach to this problem can be taken when the paths of the droplets are considered given *a priori*. This assumption leads to a scheduling problem, where the array cells en route are the limited resource that must be shared among different droplets. Griffith and Akella [37] show a solution with standard optimization tools guided by some user input, building on related work in coordinating multiple articulate robots [38], [39]. Many more references to related work in the areas of robot motion planning, flexible manufacturing systems, queuing theory and networking are also given in [37].

A related technique was used by Zhang *et al.* [5], [6] and Ding *et al.* [40], who attack the problem from the very large scale integration (VLSI) design perspective. As in [38] and [39], this approach leads to an integer programming formulation. Both groups show NP-hardness of the scheduling problem even for fixed droplet routes.

VLSI circuit routing techniques could also be employed, which address the path planning problem but do not apply directly to the inherently two-dimensional layout of the droplet-based microfluidic platform.

This paper takes a different approach, by permitting the droplet paths to be chosen freely (except for constraints defined by the microfluidics hardware). Each droplet is interpreted as a point robot moving in a discrete two-dimensional configuration space. Under this assumption, path planning of the droplets becomes a motion planning problem with multiple moving

robots. Erdmann and Lozano-Pérez showed in 1987 that this problem is NP-hard, but presented an algorithm that may find a good solution in polynomial time [41]. Their approach assigns priorities to each robot (droplet) and generates paths successively, starting with the highest priority robot. Lower priority robots consider higher priority robots as time-varying obstacles that must be avoided. The algorithm is not complete, and generated solutions depend on the priority ranking of the robots and may not be optimal.

In [42], the problem was described as a graph search, and search techniques such as A^* were suggested. Even though this brute-force approach, unlike the other work mentioned above, guarantees optimality and completeness, it is not practical for larger scale problems because of its computational complexity, which is exponential in the number of moving droplets. Böhringer [43] introduced a formal problem definition and showed initial results with a more efficient approach based on Erdmann's algorithm [41].

III. DMFS FORMAL HARDWARE SPECIFICATION

Let us briefly review the most important physical properties and design parameters of a DMFS. Motivated by these characteristics, we can then develop an abstract DMFS model that captures the essential operational features without depending on specific implementation details.

A. DMFS Design Specifications

- 1) Layout: Typically, a DMFS consists of a planar rectangular array A with $m \times n$ cells (but, e.g., an arrangement of hexagonal cells would also be possible).
- 2) Control circuitry: Various addressing schemes are possible to activate individual cells in a DMFS. In different physical implementations of DMFS, we can distinguish, for instance, individually addressable electrodes for each cell (e.g., [36]), or simpler row/column addressing (e.g., [44] and [45]). For the latter, entire rows and columns are activated, and the droplet is attracted to a neighboring cell A(x,y) only if it lies at the intersection of active column x and row y.
- 3) Parallelism: The DMFS controller may be capable of simultaneous activation of more than one cell, which will allow simultaneous motion of multiple droplets. The total number of addressable cells may be limited by a number significantly smaller than $m \times n$.
- 4) Location of cells with special functions: Droplet generators, reservoirs, cells for merging and splitting of droplets, sensors, waste, etc. may require dedicated cells with special embedded hardware. These cells may not be available when planning a droplet path across the array.

These specifications provide a physical framework within which a DMFS can operate. Based on this framework, we can establish a formal description of the problem of controlling droplets in a DMFS. Once a sufficiently general DMFS model exists, we can investigate algorithmic solutions at an abstract level without worrying about the varying details of specific hardware implementations.

B. Abstract DMFS Specification

A DMFS is specified by the droplets on the DMFS array, the DMFS hardware itself, and the task to be performed.

1) Droplets: Droplets are described by their type T and their volume V. We are assuming here that all droplets in the DMFS have the same volume, except when two droplets have been merged. Therefore, we require that a merge operation is always immediately followed by a split operation that restores the original droplet volumes.

The droplet type T is a subset of all elementary droplet types, which we describe in general as a set $\mathcal{T} = \{T_1, T_2, T_3, \ldots\}$; thus, T is an element of the power set of \mathcal{T} , $T \in \mathcal{P}(\mathcal{T})$. For example, if T_1 represents "deionized (DI) water," T_2 "methanol," and T_3 "isopropanol (IPA)," then a droplet of type $T = \{T_1, T_3\}$ describes a mixture of DI water and IPA. Note that this convention provides a simple representation of mixed droplets but does not keep track of sample concentrations. If needed, different concentrations could be represented as different elementary types.

2) DMFS Arrays and Tasks: The DMFS consists of an array A of $m \times n$ cells. Each cell in the array is either empty or occupied, which we represent by specifying its droplet type T. Thus, the DMFS can be described by $A(x,y) = T_{x,y}$ for $(x,y) \in \{1 \dots m\} \times \{1 \dots n\}$ and $T_{x,y} \subseteq \mathcal{T}$. As a special case, $T_{x,y} = \varnothing$ indicates an empty cell. We call $A \in \mathcal{P}(\mathcal{T})^{m \times n}$ the state of the DMFS.

The location of a droplet can be specified by the pair $(x, y) \in \{1 \dots m\} \times \{1 \dots n\} = C$; thus, C is the configuration space [46] of a single droplet, and C^d is the configuration space of d droplets, which we also call the droplet placement of the DMFS.

Time is assumed to be a discrete counter $t \in \{0, 1, 2, \ldots\}$, i.e., transitions in the array occur in integer time steps from t to $t + \Delta t$, where $\Delta t = 1$ unless noted otherwise. We write A_t to refer to the state of the array at a specific time t.

At this point, we can already outline the definition of a DMFS task: Given a start state $A_s \in \mathcal{P}(\mathcal{T})^{m \times n}$ and a goal state $A_g \in \mathcal{P}(\mathcal{T})^{m \times n}$, we need to find a timed sequence of valid transitions that results in the desired droplet motions from A_s to A_g . Various kinds of transitions exist; they include simple droplet transport from cell to cell, but also droplet generation, disposal, merging, and splitting. In addition to motion, droplets may also be modified by operations on cells that change their type. All these operations are chosen from the following list of valid droplet transitions, which are usually associated to specific cells or groups of cells on the array:

- 1) Droplet generation: For $(x,y) \in C$ and some $T \in \mathcal{P}(\mathcal{T})$, a droplet is generated at coordinate (x,y) if $A(x,y) = \varnothing$ at time t and A(x,y) = T at time $t + \Delta t$.
- 2) Disposing: Definition analogous to droplet generation.
- 3) Moving: Let (x,y) and $(x',y') \in C$ and |x-x'|+|y-y'|=1 (i.e., A(x,y) and A(x',y') are directly adjacent). At time t, A(x,y)=T and $A(x',y')=\varnothing$ and at time $t+\Delta t$, $A(x,y)=\varnothing$ and A(x',y')=T.
- 4) Merging: Let (x,y), (x',y'), and $(x'',y'') \in C$ such that (x',y') and (x'',y'') are directly adjacent to (x,y) but not adjacent to each other. At time t, $A(x,y) = \varnothing$,

 $A(x',y') = T_1$, and $A(x'',y'') = T_2$, and at time $t + \Delta t$, $A(x,y) = T_1 \cup T_2$ and $A(x',y') = A(x'',y'') = \varnothing$, where $T_1 \cup T_2$ is the droplet type that results in merging droplet types T_1 and T_2 .

- 5) Splitting: Definition analogous to merging.
- 6) Checking: For $(x, y) \in C$, we require that a droplet remains at A(x, y) from time t to time $t + \Delta t$. This allows, for example, sensing operations to be performed that neither change the location nor the type of the droplet (e.g., fluorescence detection).
- 7) Changing: For $(x,y) \in C$, we define a function $f: \mathcal{P}(T) \to \mathcal{P}(T)$ such that $A(x,y) = T_1$ at time t and $A(x,y) = T_2$ at time $t + \Delta t$, and $f(T_1) = T_2$. This allows transition operations that modify the droplet type but not its location (e.g., heating/cooling for PCR).
- 8) Blocking: For $(x,y) \in C$, we define a set of forbidden droplet types $\mathcal{F}_{x,y} \subseteq \mathcal{T}$ that are not allowed on A(x,y). In particular, if $\mathcal{F}_{x,y} \equiv \mathcal{T}$, then A(x,y) is blocked for all droplets.

Finally, valid placement and motion of droplets on the array is subject to the following constraints:

- 1) Placement: To avoid accidental merging of droplets, at least one empty cell is required between two occupied cells at all times, i.e., for any (x,y) and $(x',y') \in \mathbf{C}$ with $A(x,y) \neq \varnothing$ and $A(x',y') \neq \varnothing$, |x-x'| > 1 or |y-y'| > 1.
- 2) Parallel transitions: The previous constraint on placements must in particular also hold during transitions, i.e., for all pairs of droplet placements across the transition interval $[t, t+\Delta t]$ (see Fig. 2), except when merging or splitting is intended.

IV. DROPLET PATH PLANNING

This section focuses on a central task in the control of DMFS: generating efficient paths for multiple droplets that move from a given start configuration A_s to a desired goal configuration A_g . For now, we require that the types of the droplets remain unchanged during the transition from A_s to A_g (this constraint will be removed in Section V). We will first give a simple complete algorithm based on A^* search, but find that its computational complexity is very high (exponential in the number of droplets). We then present a more efficient algorithm for the DMFS motion planning problem that trades off completeness for faster execution times, while maintaining some "local" optimality guarantees.

A. Basic A* Search

This approach maintains a graph data structure to keep track of the droplet locations in the DMFS array. At any given time t, the state of the DMFS is described by A_t and identified with a node in this graph. A transition between two states A_t and $A_{t+\Delta t}$ defines a directed edge; this transition must conform with the conditions set forth in Section III-B above. Finding an optimal control strategy to transform start state A_s into goal state A_g then becomes a standard graph search problem: The shortest path between nodes A_s and A_g can be determined,

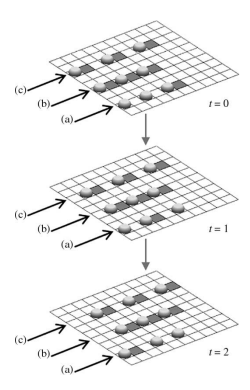


Fig. 2. Parallel droplet transitions: Droplets and their activated neighbor cells (squares) are shown at the instant when motion is commencing. The transitions in rows (a) and (c) are valid, but invalid in row (b), because during these transitions, two of the droplets have more than one activated neighbor cell, which could lead to unintentional splitting or merging.

e.g., using the A^* algorithm known from artificial intelligence programming [47].

The A^* algorithm outlined below maintains two lists of states, namely Open and Closed, which keep track of nodes that still need to be explored, and nodes that have already been processed, respectively. For each node, we maintain its predecessor p, the cost incurred g (i.e., number of transitions from A_s), the cost remaining h (i.e., number of transitions to A_g), and the total cost f=g+h. As has been widely discussed in the literature, h, which is not known in advance, can be estimated with an "admissible" heuristic function. The Manhattan metric provides such an admissible cost estimate, i.e., if droplet i at time t is at $(x_{t,i},y_{t,i})$ and its goal is $(x_{g,i},y_{g,i})$, then h(t) can be estimated as $\Sigma_i|x_{g,i}-x_{t,i}|+|y_{g,i}-y_{t,i}|$.

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Algorithm 1: A^* for droplet path planning Input: start state A_s, goal state A_g Output: shortest path from A_s to A_g Open \leftarrow \{A_s\}; Closed \leftarrow \varnothing; WHILE Open \neq \varnothing BEGIN o \leftarrow pop state with smallest f from Open; Q \leftarrow list of all valid motion transitions from o; \# Line 5 FOR each q in Q BEGIN q.g \leftarrow o.g + 1; \# q is one step beyond o q.h \leftarrow distance estimate from q to A_g; q.f \leftarrow q.g + q.h; q.p \leftarrow o; \# keep track of path from A_s via o to q IF q = A_g, RETURN q; \# goal found, success IF NOT (\exists q' \in Open such that q' = q AND q'.f < q.f)
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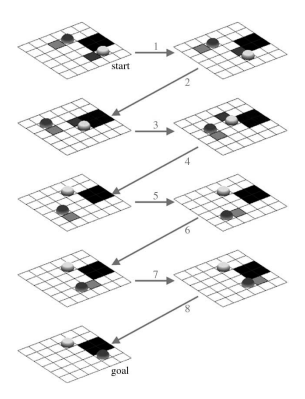


Fig. 3. Two droplets moving simultaneously on a 6×6 DMFS array while avoiding an obstacle (black cells). The two droplets start at cells (5,2) and (4,5) and trade their places in eight parallel transitions. The activated neighbor cells for their next transitions are also shown.

AND NOT $(\exists q' \in \text{Closed such that } q' = q \text{ AND } q'.f < q.f)$ THEN add q to Open; // found new state q to be

END

add o to Closed; // finished exploring node o

explored

END

RETURN Ø; // search exhausted, failure

Fig. 3 shows a simple example where two droplets swap their position while avoiding an obstacle. The A^* algorithm is guaranteed to always find an optimal solution if one exists, and indicate failure otherwise. However, the downside of this approach is its high asymptotic complexity. Suppose the number of droplets is d. In the simplest case, all are of the same type T_0 . Then the number of different placements of droplets on the array is $\binom{mn}{d}$, which for modest numbers m=n=10 and d=10 yields more than 1.7×10^{13} possibilities. If all droplets are of distinct type T_1, \ldots, T_d , this number increases by d! (to $\approx 6.3 \times 10^{19}$). One might hope that, in practice, most of these choices need not be explored. However, at each step, d droplets offer up to 4d choices to be moved, assuming four neighbor cells per droplet. Thus, finding a strategy with s steps could mean checking up to $(4^d)^s$ choices or risk missing the solution, resulting again in astronomical numbers even for s < 10.

We conclude that the search graph explored with the A^* algorithm has $\mathrm{O}((mn)!)$ nodes and a branching factor of $\mathrm{O}(4d)$, leading to a runtime complexity exponential in d, which is prohibitive for any nontrivial array size with more than a few droplets.

B. Prioritized A* Search

The discussion above has shown that droplet path planning for DMFS has two main aspects, namely: 1) generating efficient droplet path plans; and 2) finding efficient algorithms to generate these plans. Erdmann and Lozano-Pérez's [41] NP-hardness results for coordinating multiple moving objects indicate that compromises need to be made to obtain practical solutions, and completeness or optimality in motion plans has to be traded off with efficiency in plan generation. They propose to impose a priority order on the moving objects, and sequentially find "locally" optimal solutions. In our case, the order can be assigned at random, or based on application-specific guidelines (e.g., water may have lower priority than droplets containing expensive or volatile compounds):

Algorithm 2: Prioritized A^* for droplet path planning Input: start state A_s , goal state A_g , priority order for droplets Output: path from A_s to A_g

 $S \leftarrow \varnothing$; // partial prioritized solution

FOR all droplets i in decreasing priority order BEGIN

call Algorithm 1 to determine an optimal path for droplet i while considering all droplets with higher priority as moving obstacles and ignoring all droplets with lower priority;

IF solution for droplet i exists THEN add solution to S; ELSE RETURN \varnothing ; // failure

END

RETURN S; // success

Fig. 1 was generated using this algorithm. It eliminates the exponential complexity in d, where d is the number of droplets in the DMFS. Instead, the prioritized algorithm is linear in d. As a tradeoff: 1) it is no longer complete—existing solutions may be missed; and 2) the solution may not be "globally" optimal while each droplet i finds a "locally" optimal path among the moving droplets of higher priority, the complete solution will in general depend on the priority order and not be "globally" optimal. Thus, as was pointed out in [41], selecting the priority order can greatly influence the final solution. For instance, if a short path is important for a specific droplet type, then it should receive high priority. However, total runtime is dominated likely by low priority droplets, since they may take convoluted paths to circumnavigate all higher priority droplets. A good heuristic for assigning priorities will take these points into account, as well as other application-specific factors. For example, droplets whose type appears frequently on the DMFS could be assigned lower priorities than rare droplet types, because it is likely that one of the abundant droplets is already close to a desired destination.

C. Parallel Droplet Motion

The algorithms given so far are able to generate plans with simultaneous motion of multiple droplets. Beside the physical limitations to parallelism discussed with Fig. 2, the DMFS control hardware may impose additional constraints. For example, [44] and [45] describe a DMFS with simpler row/column

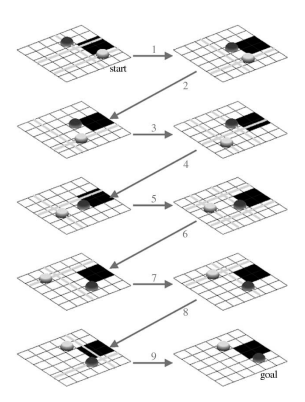


Fig. 4. Two droplets trading places as in Fig. 3, but here droplets move only to neighbor cells whose row and column have been activated (indicated by a line). An optimal strategy now requires nine steps. Note that even though parallel droplet motion occurs in several steps, transitions 1 and 4 in Fig. 3 would not be possible with this addressing scheme.

addressing, where a droplet moves to a neighboring cell A(x,y) only if it lies at the intersection of activated column x and row y. Such conditions are encoded in line 5 of Algorithm 1:

" $Q \leftarrow$ list of all valid motion transitions from o;"

Generation of this list of transitions must be implemented depending on the hardware specifications. Fig. 4 shows an optimal solution for the same start and goal states as in Fig. 3 but with this more limited row/column addressing scheme.

D. Duplicate Droplet Types

An important special case occurs when multiple droplets in the DMFS have the same droplet type. This is a likely scenario in practice, especially in DMFS with large numbers of droplets. In this case, there is no unique mapping between droplets in A_s and A_g (or with any intermediate state A_t). This complicates the calculation of the cost estimate h, but can also provide for more efficient plans by choosing opportune droplets that are closest to their respective goals.

Suppose we are given two sets of d droplet placements, S_1 and $S_2 \in \mathbb{C}^d$, and all droplets have the same type T. We can find the minimum cost match between S_1 and S_2 efficiently (in analogy to bipartite graph matching [48]) by a greedy algorithm that sequentially matches up coordinate pairs with minimal Manhattan distance until all coordinates are paired up. This pairing leads to a monotone underestimate of the actual cost, and can, thus, be used as an admissible estimate for h. With this addition, both the basic and the prioritized A^* algorithm for droplet path planning can efficiently handle inputs with duplicate droplet types.

V. DMFS TASK PLANNING

The previous section addressed the DMFS motion planning problem. However, transitions of the droplet types (due to mixing or other processing as discussed in Section III) are essential parts of DMFS operation. Thus, this section extends the previously introduced algorithms to the general DMFS planning problem, which allows all remaining droplet transitions listed in Section III-B, including merging, splitting, and changing of droplet type. This ultimately leads to the much broader question of how to transform a general laboratory protocol into a specific sequence of commands that can be executed on a DMFS.

A. Basic Graph Search

A straightforward algorithm to solve the general DMFS planning problem can be derived from Algorithm 1, where we can again modify line 5 to allow the complete set of transitions listed in Section III-B, also including in particular changes of droplet type. However, this causes some immediate problems: 1) the number of possible transitions from each state (i.e., the branching factor in the search graph) becomes very large; and 2) it is difficult to find an admissible heuristic for the A^* algorithm, causing it to degenerate into breadth-first-search; in combination, this would result in very inefficient searches. These problems would apply equally to a modified Algorithm 2.

B. DMFS Task Protocols

To develop a more useful algorithm, it is important to keep in mind that the tasks to be executed here typically are laboratory protocols. Thus, it is reasonable to assume that the user (e.g., a chemical engineer or a researcher in molecular biology) has carefully worked out the individual steps in this protocol and identified the intermediate products that are being generated during its execution. With this additional input, we can find efficient algorithms to perform these tasks on a DMFS, while leaving the task design to a knowledgeable human operator.

We now introduce a very simple DMFS task language; the user of a DMFS specifies the tasks to be executed in this language, based on the laboratory protocol for the process of interest. Our Algorithms 3 and 4 then interpret this task description and translate it into actual DMFS commands.

DMFS Task Language // Textual description of DMFS protocol // $x \in \{1 \dots m\}; y \in \{1 \dots n\}; \Delta x, \Delta y \in \{0, 1, \dots\}; t \in \{1, 2, \dots\}$ // $T \in \mathcal{P}(T); f : \mathcal{P}(T) \to \mathcal{P}(T)$ // id is an arbitrary textual identifier for a cell in x y T id [time t] out x y T id [time t] waste x y id mergesplit $x y \Delta x \Delta y T$ id [time t] check x y T id [time t] change x y f id [time t] block x y T id connect from-id to-id

The statements in this language correspond to the DMFS array transitions listed in Section III-B with the following additional explanations:

- 1) x and y are the cell coordinates in the array. In general, we assume that transitions happen on a single array cell, except merge/split operations, which may require larger cells (specified by Δx and Δy such that $x + \Delta x \leq m$ and $y + \Delta y \leq n$).
- 2) A droplet of type T_d is allowed on a cell with specified type T only if $T_d \subseteq T$.
- 3) in, out, mergesplit, check, and change have an optional argument time t with default value t=1 that specifies the time required for the transition.
- 4) "waste $x \ y$ id" is a short form for "out $x \ y \ T$ id" which implies that the droplet type does not matter because the droplet will be discarded.
- 5) "block x y T id" prohibits any droplet of type $T_d \subseteq T$.
- 6) "**connect** from-id to-id" implies a single droplet moving between the two specified cells.

Note that identifiers need not be unique. However, if multiple transitions have the same identifier, then they belong to the same cell group and must describe the same transition. For example, we can write "in 1 1 H₂O DI input" and "in 3 1 H₂O DI input" to specify two cells (1, 1) and (3, 1) that provide a supply of DI water. Thus, if we write "connect DI input mix" then our algorithm will choose one of the DI water inputs to route a droplet to the cell with identifier "mix."

Fig. 5 gives a sample DMFS task input. Four input droplets of three different initial droplet types go through a sequence of merges, splits, type transitions, and checks, before finally reaching an output or waste cell. The user specifies these steps and their locations on the DMFS array. Our algorithms automatically generate the order of these operations, the selection of specific cells from cell groups, and the exact droplet paths and schedule.

In large DMFS with many moving droplets and many in, out, mergesplit, check, and change cells, choosing the locations where droplets are processed should also be automated. A greedy algorithm and simulated annealing are discussed in [49] to attack this NP-hard layout problem.

While this list of statements may look tedious, it is simply a textual description of a graph in which every node represents a transition (**in**, **out**, **waste**, **mergesplit**, **check**, **change**) at a specific location on the DMFS, and every edge corresponds to a droplet motion (**connect**). We call this directed graph, which specifies the flow of the droplets through the DMFS, the task graph. It gives a more intuitive representation of the DMFS task to be executed and will be discussed in the following section.

C. DMFS Planning Algorithm

The final part of this paper is dedicated toward translating a DMFS task description, given in the language from the previous subsection, into a sequence of commands that can be executed on the array. This algorithm will do the following: 1) Generate the task graph from the textual input. 2) Identify initial transitions (typically, **in** nodes) that do not have any incoming edges. 3) Assign levels to all nodes in the task graph

```
in 00 {R} red
in 0.2 \{G\} green
in 0 4 {B} blue
in 0 6 {B} blue
mergesplit 4 2 0 0 magenta
   connect red magenta
   connect blue magenta
mergesplit 4 5 0 0 cyan
   connect green cyan
   connect blue cyan
mergesplit 4 8 0 0 pale
   connect magenta pale
   connect cvan pale
change 14 1 \{R,B\} \rightarrow \{M\} modify
   connect magenta modify
change 14 3 \{G,B\} \rightarrow \{C\} change
   connect cyan change
change 14 5 {R,G,B}→{W} process
   connect pale process
check 10 2 all sensor
check 10 5 all sensor
   connect modify sensor
   connect change sensor
out 20 0 all out
out 20 2 all out
out 20 4 all out
   connect process out
   connect pale out
waste 20 10 trash
   connect sensor trash
   connect sensor trash
```

Fig. 5. Sample DMFS tasks. There are three cell groups, namely blue, sensor, and out, consisting of multiple cells with the same identifier and the same transitions (**in**, **check**, and **out**, respectively). The keyword **all** indicates the entire set of droplet types \mathcal{T} . Note that only two out of the three **out** cells will be used.

according to their precedence relationships such that transitions on the same level can be executed in parallel.

```
Algorithm 3: Task graph generation
Input: DMFS task description tasks
Output: task graph G with level assignments
parse tasks and generate the corresponding task graph G;
old \leftarrow \varnothing;
new \leftarrow all \text{ nodes in } G;
current \leftarrow all nodes in G that do not have predecessors;
   WHILE current \neq \varnothing BEGIN
  mark all nodes in current with level i;
  add all nodes in current to old;
  current \leftarrow all nodes in new that have only predecessors
  in old:
  remove all nodes in current from new;
  i \leftarrow i + 1;
END
IF new = \emptyset
THEN RETURN G (with level numbers); // success
```

If the directed graph is acyclic, then this algorithm finds a level assignment with a minimum number of levels (which we call l), thus, maximizing the potential for parallel execution of the transitions represented by its nodes and edges. Note that

ELSE RETURN ∅; // failure

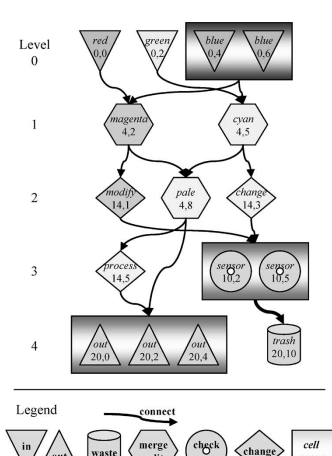


Fig. 6. Task graph with level assignments generated from the task description in Fig. 5. Transitions on the same level can be executed in parallel. (a) There are two droplets moving from the sensor cell group to trash, indicated by a double thickness arrow. (b) Only two of the three output cells will be used.

these level assignments merely reflect precedence relationships, not actual execution times: Droplet transitions on a specific level and droplet motions between levels may have varying transition times (and the latter are not yet known). Thus, faster droplets may have to wait until slower droplets are finished on each level.

Algorithm 3 assumes that there are no resource conflicts between droplets on any given level, i.e., no two transitions require the same cell on the DMFS array. If this cannot be guaranteed during the specification of the DMFS task, then the algorithm must be modified to assign conflicting transitions to different levels. See, e.g., [50] for a comprehensive approach to dealing with such resource constraints.

Fig. 6 shows the task graph generated by Algorithm 3 from the DMFS command input given in Fig. 5. From its level assignment, we can immediately generate array states A_{i-} and A_{i+} that correspond to each level $i \in \{0,\ldots,l\}$, such that A_{i-} and A_{i+} are the state of A immediately before and after the transitions of level i, respectively. Then, we can use Algorithm 2 to determine the droplet motions between arrays A_{i-1+} and A_{i-} for all $0 < i \le l$:

Algorithm 4: DMFS planning Input: DMFS task description *tasks*

Output: task graph G and corresponding droplet motions S

TABLE I DMFS TASK STATE AND TRANSITIONS

State A ₀ +	Droplet Placements			
	(0,0) {R}	(0,2) {G}	(0,4) {B}	(0,6) {B}
	\downarrow	\downarrow	\downarrow	\downarrow
A ₁ - +	(4,2) {R} {R,B}	(4,5) {G} {G,B}	(4,2) {B} {R,B}	(4,5) {B} {G,B}
	\downarrow	↓	↓	\downarrow
A ₂ - +	(14,1) {R,B} {M}	(14,3) {G,B} {C}	(4,8) {R,B} {R,G,B}	(4,8) {G,B} {R,G,B}
	\downarrow	↓	↓	•
A ₃ - +	(10,2) {M} {M}	(10,5) {C} {C}	(14,5) {R,G,B} {P}	(4,8) (R,G,B) {R,G,B}
	\downarrow	↓	\downarrow	\downarrow
A ₄ _	(20,10) {M}	(20,10) {C}	(20,4) {P}	(20,2) {R,G,B}

Start, intermediate, and goal states generated from the task graph in Fig. 6. For each state A_i , $i \in \{0, \dots, 4\}$, droplet placements and their respective types are shown before and after the transition on level i. Three or four droplets move simultaneously during the four transitions from A_{i-1} to A_{i-1} (for i > 0), indicated by downward arrows.

 $G \leftarrow \text{call Algorithm 3}$ with input tasks

IF $G = \emptyset$ THEN RETURN \emptyset ; // no task graph exists, failure $S \leftarrow \emptyset$;

FOR $i \leftarrow \{1 \dots l\}$ BEGIN // l is the maximum level number of G

determine A_{i-1+} and A_{i-} , using G and S;

 $S_i \leftarrow$ call Algorithm 2 with start A_{i-1+} and goal A_{i-} ; IF $S_i = \emptyset$

THEN RETURN \emptyset ; // no droplet path i exists, failure ELSE add S_i to S; // droplet path i found

END

group

RETURN G and S; // success

Table I lists all the states and transitions generated by Algorithm 4 from the task graph in Fig. 6. Fig. 7 attempts to visualize parallel motion of multiple droplets on the DMFS for the transition from A_{3+} to A_{4-} .

Algorithm 3 is linear in the number of nodes and edges in the task graph. The complexity of Algorithm 4 is dominated by the calls to Algorithm 2, which occur l-1 times total. These algorithms were implemented in Java. The total runtimes for the examples in this paper are in the millisecond range. The code is available upon request from the author.

VI. CONCLUSION

This paper makes the following contributions: 1) a formal hardware-independent model of DMFS; 2) novel algorithms for motion and task planning with DMFS, leading to efficient (albeit not necessarily optimal or complete) solutions for

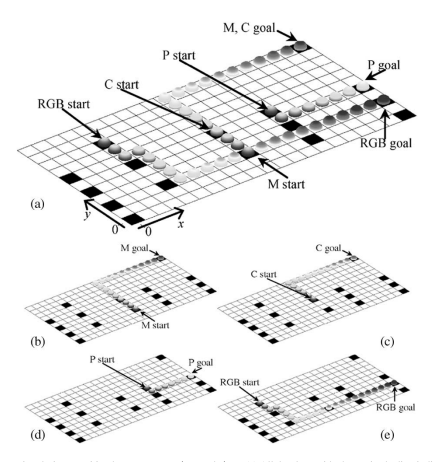


Fig. 7. Simultaneous droplet motion during transition between states A_{3+} and A_{4-} . (a) All droplets, with change in shading indicating progressing time. Cells with special functions are marked as black squares. (b)–(e) Individual droplet paths for the droplets of type $\{M\}$, $\{C\}$, $\{P\}$, and $\{R,G,B\}$, respectively. In (b) and (c), droplet $\{C\}$ follows the path of droplet $\{M\}$ at a distance of three cells; in (e), the droplet circumnavigates the mergesplit cells at (4,2) and (4,5) but is allowed to pass over the sensor cell at (10,2).

coordinating large numbers of simultaneously moving droplets on a two-dimensional array; 3) an approach to automate the transition from general laboratory protocols to DMFS control command sequences; and 4) results using an implementation of these algorithms in Java.

The developed models and algorithms are "modular," such that results from the different sections are largely independent; e.g., DMFS task planning in Section V does not rely on a particular droplet path planning algorithm so some other algorithm could be readily substituted for prioritized A^* . Similarly, the path planning algorithms from Section IV could be applied to a different task planning algorithm.

Droplet manipulation based on electrowetting on arrays with up to hundred cells has been demonstrated by several groups (e.g., [3], [44], and [51]), and an electrophoresis-based system with integrated CMOS addressing of tens of thousands of cells by [34]. The computational complexity for generating optimal droplet motion plans has been shown to be prohibitive even for much smaller systems. Thus, we have focused on finding an acceptable tradeoff between efficiency and optimality.

A very different approach to this problem could be to limit droplet manipulation to a few standard "prepackaged" strategies. For example, on a 100×100 array, about 50 droplets could move in parallel across the array, followed by another wave of 50 droplets, etc., resembling a repetitive "peristaltic" motion [43]. However, in this case, the fundamental advan-

tage of flexibility and reprogrammability in DMFS versus conventional (channel, valve, and pump based) microfluidic architectures is lost. In addition, the question still remains how to initially generate the "prepackaged" strategies if they involve more complicated motion paths by many simultaneously moving droplets.

Other future work should explore the following directions.

- 1) Polynomial approximation algorithms exist for NP-hard problems (e.g., traveling salesman [52], [53]), which guarantee a tight limit on nonoptimality. If, e.g., a control strategy for a complex DMFS can be generated in polynomial time that is guaranteed to be at most twice as long as an optimal solution then this might be sufficient for most practical purposes.
- 2) While the (prioritized) A^* algorithm has been effective in solving graph search problems, it is incomplete and worst case exponential in the branching factor. More detailed benchmark tests could provide insights about scenarios where the algorithm fails to find solutions efficiently.
- 3) The optimal level number l produced by Algorithm 3 does not automatically imply maximal parallelism in droplet motion. Some nodes in the task graph can be assigned to a range of levels without affecting l, but varying level assignments may produce droplet motion plans with varying efficiency. For example, the droplet motion from

- (4,8) to (20,2) in Table I and Fig. 7(e) can be executed during transition $A_{2+} \rightarrow A_{3-}$ or during $A_{3+} \rightarrow A_{4-}$.
- 4) A related question is whether it is essential to allow parallel droplet motion in line 5 of Algorithm 1. An alternative approach would first generate plans without parallelism, and then postprocess the generated plan to identify all droplet motions that could be executed in parallel.
- 5) More generally, it may be possible to improve the output of Algorithm 2 with some postprocessing that locally improves the droplet motions.
- 6) The previous three points hint that our DMFS formalism could be developed much further. A general approach in this direction based on state complexes was given recently in [54], which presents efficient algorithms to detect and optimize parallelism.
- 7) As mentioned in Section III-A, parallelism may be limited by the hardware controller to a number smaller than the total droplet count. This was not explicitly addressed in this paper, but could again appear as an additional constraint in line 5 of Algorithm 1.

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