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Evaluation of Telerobotic Shared Control Strategy for Efficient Single-Cell Manipulation

Jungskik Kim, Hamid Ladjal, David Folio, Antoine Ferreira, Member, IEEE, and Jung Kim, Member, IEEE

Abstract—Microinjection using a glass capillary is a highly efficient method for the delivery of exogenous materials into cells and is widely used in biomedical research areas such as transgenics and genomics. However, this direct injection is a time-consuming and laborious task, resulting in low throughput and poor reproducibility. Here, we describe a telerobotic shared control framework for microinjection with high manipulation efficiencies, in which a micromanipulator is controlled by the shared motion commands of both the human operator and the autonomous controller. To determine the optimal gains between the operator and the controller, we proposed a quantitative evaluation method using a model of speed/accuracy trade-offs in human movement. The results showed that a 40–60 % weighting on the human operator (or the controller) produced the best performance for both speed and accuracy of task completion suggesting that some level of both automation and human involvement is important for microinjection tasks.

Note to Practitioners—In single-cell microinjection, for the small size and delicate structure of a cell, to date, most human operators have manipulated biological cells manually; therefore, low manipulation efficiency and poor reproducibility has been reported for this task. Most manipulation systems have primarily focused on limited visual feedback in conjunction with a dial-based console system, requiring extensive operator training to perform injection tasks with reproducible results. To address these problems, a telerobotic shared control method for microinjection was developed by integrating the automatic and direct manipulation functions of a robotic system. While a controller retains cells and glass pipettes within a desired path or space, the operator can concentrate on the injection task, thus achieving high throughput and dexterity.

Index Terms—Fitts’ and steering laws, microrobotic control, single-cell microinjection, telerobotic shared control.

I. INTRODUCTION

The highly efficient transfer of foreign materials into cells remains a challenge in biotechnology, both for fundamental cellular and molecular biology research and in biomedicine. Several methods have been developed for the successful delivery of exogenous materials into cells. Among them, single-cell microinjection is performed to directly introduce foreign materials, such as DNA, proteins, sperm and drugs, into individual cells [1]-[3], and it is a relatively more efficient delivery method than electrical [4], viral [1], chemical [1], and other transfer methods [5], [6]. Microinjection is used in many different biotechnologies such as cell transfection with DNA in mammalian cells [1], induced pluripotent stem (iPS) cell transfer [7], and in vitro fertilization (IVF) by intracytoplasmic sperm injection (ICSI) [8], [9]. However, although microinjection has relatively high delivery efficiencies, the injection task is time-consuming and labor-intensive work that limits the manipulation of large numbers of single cells [1], [2]. The elaborate manipulation of small, delicate cells demands certain technical requirements, such as the development of micro-motion stages and the fabrication of micro-pipettes with a fine tip. In addition, great manipulation skills are required of a human operator requiring extensive training to perform these injection tasks; thus, microinjection has low manipulation efficiencies, resulting in low throughput and poor reproducibility.

The manipulation of micrometer-sized cells is currently hindered by the limited capability of sensing technologies. These manipulations are often performed with insufficient visual information, within a small field of view and a narrow depth of field, creating a difficult situation for the detection and positioning of cells or glass pipettes. In addition, there is no force/tactile feedback for robot control or haptic interaction. Insensate glass pipettes can easily damage themselves or the target cells. To address these problems, several single-cell microinjection systems have been proposed to improve the manipulation efficiency. Automated microinjection systems have been developed to remove human involvement from the injection process [10]-[16], where a visual servoing approach is usually used to control the position and force of a micromanipulator; however, it is challenging to create a fully automated manipulation system because microinjection is conducted under diverse and complex conditions such as varying cell size (from one micrometer to hundreds of micrometers), cell types (e.g., suspended or attached ones) and liquid environments. Fig. 1 compares the various cells typically manipulated in many research and clinical fields, showing that the differences in the geometry and the material properties are greater than one-thousand-fold and one-million-fold, respectively. Therefore, there are difficulties in the dexterous manipulation of cells with multiple degrees of freedom (DOF) and in target selection (e.g., cell nucleus or cytoplasm) in visual servoing [17], [18]-[20]. Teleoperated microinjection systems have been developed to provide haptic feedback during manipulation.
[21]-[25]. However, most of them have provided force sensing and feedback for only a small number of DOF.

Here, we present a telerobotic shared control (TSC) framework developed for single-cell microinjection with high manipulation efficiency. The motivation of the TSC arose from the idea that the collaboration of a human and a robotic system can increase the quality and capability of manipulation by exploiting a human’s ability to skillfully manipulate objects with dexterity and disturbance adaptation along with a robot’s accuracy and repeatability [26]-[28]. To reduce the difficulties in biomimicromanipulation mentioned above and simultaneously achieve high throughput and dexterity, both automatic and direct manipulation functions of the system are needed in microinjection. In the TSC approach, a human operator can control the manipulator as much as possible while a controller retains cells and glass pipettes within a desired manipulation path or space to provide adequate performance. In the remainder of this paper, the TSC strategy is first presented, and the sharing method for the teleoperation and automation modules of the microinjection system is described. In addition, we provide a quantitative analysis to determine what level of automation (or direct manipulation) is needed for optimal TSC. Most previous studies on shared control have not addressed how to determine the degree of autonomy (or human involvement) to implement in the telemanipulation for the best performance. Although shared control has been applied to various applications [28]-[31], this work is, to the authors’ knowledge, the first providing an evaluation method for determining the optimal shared control gains and the first application of shared control in a cellular micromanipulation. This work focused on the evaluation of a shared control for microinjection using the proposed model with Fitts’ and steering laws.

II. Method

A. Telerobotic Microinjection System Design

A microinjection task consists of guiding a micropipette to a cell, puncturing the cellular membrane, and depositing foreign materials at a targeted position. The cells in the injection chamber with a liquid medium are attached to a substrate or suspended by a second (holding) glass pipette. The pipette tip is angled to approach a cell in a liquid medium.

The telerobotic biomimicromanipulation system for microinjection consists of a master robot to input the operator’s motion command, micromanipulators with glass micropipettes and a vision system (Fig. 2).

The task space of the master robot is represented by the coordinate frame \( \{M : (x,y,z)\} \in \mathbb{R}^3 \), the task space of the micro manipulator is represented by the coordinate frame \( \{S : (X,Y,Z)\} \in \mathbb{R}^3 \), and the image space of the fixed camera is defined by the image coordinate frame \( \{I : (u,v)\} \in \mathbb{R}^2 \).

Visual information is a main sensing modality in the biomimicromanipulation tasks. The processing of visual data determines the focal plane and the guidance path of the pipette tip within the image frame. In this study, because the input of the manipulator is given in the slave frame \( \{S\} \), the mapping between \( \{S\} \) and the image frame \( \{I\} \) forms a critical component for servoing the pipette tip. To compute this mapping, a calibration methodology was developed and implemented. Several calibration methods exist in the literature and are mostly used in macroscale vision applications [47]. Unfortunately, those methods cannot be directly applied to calibrate an optical microscope due to the characteristics...
of the optical system (e.g., high optical magnification and very narrow depth of field), and some dedicated methods have, therefore, been proposed [48]. In this work, the camera calibration parameters are defined in the focal plane (image space); therefore, the non-sharp tip image, due to the depth directional motion of the robot, is ignored. For an orthographic camera projection model, the tip position $\mathbf{p} = (X, Y, Z)^T$ in the slave frame can be related to its position $\mathbf{p'}$ in an image space as follows:

$$\mathbf{p'} = \left( \frac{u}{v} \right) = \left( \frac{X}{S_X + u_0} \frac{Y}{S_Y + v_0} \right) \quad (1)$$

where $S_X$ and $S_Y$ are the pixel dimensions ($\mu$m/px) and the initial tip position $\mathbf{p}_0 = (u_0, v_0)^T$ is measured by the template-matching method in the focal plane [25].

When assuming an accurately position-controlled micromanipulator and an orthographic projection model, it is important to initially locate the tip in the focal plane. In this work, the precise placement of the tip along the $Z$ depth is achieved by the comparison of the pixel gradient magnitudes at the tip because the tip is angled and has pure translational motion. The gradient magnitudes are computed along the center line in the $u$ direction of the tip, and the maximum values for each $Z$ depth are compared. The gradient magnitude is at a maximum when the tip is in the focal plane (Fig. 3).

**B. Telerobotic Shared Control**

The overall control architecture of the TSC system is described in Fig. 4. The TSC has two levels: the lower level consists of the direct manipulation and the autonomous controller based on an artificial potential field, and the higher level is the shared controller to integrate both modules.

The total input command $\mathbf{s}\mathbf{u}$ for the micromanipulator is defined as the weighted sum of motion commands from the operator ($\mathbf{u}_h$) and the autonomous controller ($\mathbf{u}_c$):

$$\mathbf{s}\mathbf{u} = K_h\mathbf{u}_h + K_c\mathbf{u}_c$$

$$K_h + K_c = 1$$

$$0 \leq (K_h, K_c) \leq 1 \quad (2)$$

where $K_h$ and $K_c$ are the weighting factors for the operator and the controller, respectively.

The input from the human operator is the position command of the micromanipulator under position control: $\mathbf{u}_h = \alpha_p \mathbf{M}\mathbf{p}$, where $\mathbf{M}\mathbf{p}$ is the Cartesian tip position of a master device and $\alpha_p$ is the scaling factor. An artificial potential field-based control algorithm [49] is applied for the autonomous manipulation module, in which the manipulator is considered to be a particle and is controlled under a force field by attractive and repulsive potential functions. An attractive potential field is constructed to attract the manipulator to the guidance path, and a repulsive potential field is generated around a cell membrane to prevent the micro pipette tip from passing the membrane in any direction other than the injection direction. The total potential field is defined by the sum of both potential fields, and each potential function is defined as follows:

$$U_{total}(\mathbf{p}) = U_{att}(\mathbf{p}) + U_{rep}(\mathbf{p})$$

$$U_{att}(\mathbf{p}) = \frac{1}{2} K_d | \mathbf{p}_d - \mathbf{p} |^2$$

$$U_{rep}(\mathbf{p}) = \frac{1}{2} K_r | \mathbf{p} - \mathbf{p}_0 |^2, \quad \text{for} \ | \mathbf{p} - \mathbf{p}_0 | \leq r \quad (3)$$

where

$\mathbf{p}_d =$ nearest point from the tip to the guidance path

$\mathbf{p}_0 =$ nearest point from the tip position to the membrane

$K_d, K_r =$ constant gains

$r =$ radius of influence.

Fig. 5 shows a two-dimensional example of a potential map for a zebrafish embryo. Finally, for the net force $\mathbf{f}_p$ is defined by the negative gradient of the potential field, and the input command from the artificial potential field algorithm is defined as follows:

$$\mathbf{u}_c = c_f f_p = -c_f \nabla U_{total} \quad (4)$$

where $c_f$ is the compliance constant for modifying the input command in $\mu$m/N.

**C. Optimal Shared Control Gain Tuning**

In the TSC, the input commands from the teleoperation module and the autonomous module to the shared controller
have gains (weighting factors) of $K_h$ and $K_c$, respectively. These gains play an important role in deciding which module will be weighted more in the TSC. High human operator weight can lead to the microinjection system acting as a conventional direct manipulation system. Conversely, when more weight is given to the automation module, the human operator can lose control of the micromanipulator.

In this work, to quantitatively evaluate the effect of the weighting level of each module on the TSC and so obtain the optimal gains, we performed 2D pointing and steering tasks based on Fitts’ law and the steering law. Fitts’ law and the steering law were first developed to quantify the speed and accuracy trade-off in target-directed movements [50], [51]. They have been applied in various applications, such as human-computer interaction and robotics [52]-[56].

In Fitts’ law, the Movement Time ($MT$) in seconds to select a target of width $W$ and at distance $D$ is given by

$$MT = a + b \log_2 \left( \frac{D}{W} + c \right)$$

where $a$ and $b$ are empirical parameters determined by linear regression, and $c$ is 0, 0.5 or 1 (we selected 1, following Mackenzie and Buxton [57]). The term $\log_2(D/W + c)$ refers to the Index of Difficulty ($ID$) in bits, which represents the difficulties of the tasks.

In a typical Fitts’ law formulation, the target-pointing movement is only considered between the initial and final positions, and it is thus not appropriate for trajectory-based tasks. Therefore, if the movement is constrained along a predefined path, the steering law accurately predicts the $MT$ with path length $D$ and width $W$ [51]:

$$MT = a + b \frac{D}{W}$$

Although shared control between humans and robots has been widely studied and various applications for Fitts’ law have been presented, the quantitative analysis for choosing ideal weighting in shared control using Fitts’ law has not been studied. Here, we present a method for determining the ideal weights for shared control using Fitts’ and steering law tests.

In the proposed TSC for microinjection, a micropipette-guiding task, i.e., directing the tip to a cell along the guidance path with depth directional motion compensation, can be seen as a steering law task, and target selection from the guidance path can be a Fitts’ law task of compensating for target-detection error. We propose a model combining Fitts’ law and the steering law, in which we hypothesized that the total index of difficulty $ID_t$ (bits) is obtained by:

$$MT = a + b ID_t$$

$$ID_t = ID_f + d \cdot ID_s$$

where $ID_f = \log_2(D/W + c)$ in (5) and $ID_s = D/W$ in (6), and $d$ is 1·bits for unit conversion.

### III. Experiment

#### A. System Setup

A biomicromanipulation workstation is shown in Fig. 6. Zebrafish embryos were used as the manipulated suspended cells. The operator commanded the master robot (SensAble Technologies, PHANToM Premium 1.0, USA) to input the motion command for the micromanipulator in the slave space. The slave consists of a micromanipulator (Physik Instrumente, F-131, Germany) with an injection needle, and the other micromanipulator (Sutter Instrument, MP225, USA) consists of a holding glass pipette. The micromanipulator has three translational motions in the X, Y, and Z direction for input...
The vision system includes a CCD camera (SVS-Vistek, SVS340MC, Germany with a 640×480 pixel resolution and maximum frame rate of 250 fps) with a microscopic lens (Moritex, MML2-ST65D with 2× magnification) to capture images for the cells and the motion of the slave tip. The position of the cell in the image space is extracted using the Hough transform as the center point of the detected circle, and the focusing of the nonangled holding pipette with a cell was implemented based on the template-matching method [10]. Finally, the guidance path is determined as the linear path from the cell position to the injection starting position near the cell membrane, as shown in Fig. 7. The gains for artificial potential field were set to $K_u = 1.0$, $K_r = 1.0$, and $c_f = 1.0$. The microinjection process is shown in Fig. 8.

B. Fitts’ and Steering Laws Test Design

Five participants (four males and one female aged 26.8 ± 2.2 years) participated in the experimental task. All participants were right-hand dominant and inexperienced in microinjection and teleoperation systems.

Fig. 9 shows the task window for the proposed model (7). The steering task was performed first for the tunnel (Box 1 in Fig. 9), in which the desired path was on the center line along the horizontal direction. The participants were asked to guide a white plus sign (the micropipette tip position in image space) from the right side to the left side of the tunnel without crossing the top or bottom sides of the tunnel. The starting position of a white plus sign was randomly chosen every trial. After passing the tunnel, a square box appeared (Box 2), and the participants were instructed to tap inside Box 2 by clicking the button on the stylus of the haptic device. If the participants failed to perform the task by leaving the tunnel or mistargeting the square box, the white plus sign changed
TABLE I

<table>
<thead>
<tr>
<th>W (px)</th>
<th>D₀ (px)</th>
<th>D₁ (px)</th>
<th>IDₜ (bits)</th>
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<tr>
<td>40</td>
<td>50</td>
<td>90</td>
<td>2.95</td>
</tr>
<tr>
<td>30</td>
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<td>70</td>
<td>6.46</td>
</tr>
<tr>
<td>30</td>
<td>150</td>
<td>80</td>
<td>6.87</td>
</tr>
<tr>
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<td>30</td>
<td>180</td>
<td>90</td>
<td>8.00</td>
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<td>90</td>
<td>90</td>
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<tr>
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<td>50</td>
<td>14.58</td>
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TABLE II

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<th>Model</th>
<th>r²</th>
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<td>0.2</td>
<td>MT = 0.6288 + 0.1133IDₜ</td>
<td>0.9317</td>
</tr>
<tr>
<td>0.4</td>
<td>MT = 0.2837 + 0.1283IDₜ</td>
<td>0.8956</td>
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<tr>
<td>0.6</td>
<td>MT = 0.1786 + 0.1485IDₜ</td>
<td>0.9030</td>
</tr>
<tr>
<td>0.8</td>
<td>MT = −0.1224 + 0.2207IDₜ</td>
<td>0.8183</td>
</tr>
<tr>
<td>1.0</td>
<td>MT = −0.2864 + 0.2698IDₜ</td>
<td>0.9138</td>
</tr>
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TABLE III

<table>
<thead>
<tr>
<th>SS</th>
<th>df</th>
<th>MS</th>
<th>F</th>
<th>* * p &lt; 0.001</th>
<th>η² = 0.7718</th>
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<td>1.2876</td>
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<td>-</td>
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<tr>
<td>Error</td>
<td>1.5232</td>
<td>16</td>
<td>0.0952</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Total</td>
<td>11.9447</td>
<td>24</td>
<td></td>
<td>-</td>
<td>-</td>
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</tbody>
</table>

C. Results and Discussion

The experimental results along with the linear regression for the proposed model (7) are shown in Table II and Fig. 10, respectively. The experimental data fitted the model with an r² value over 0.8183, showing that the MTs were predicted well by the proposed model and that the model can thus be used for the performance measurement to evaluate the system and determine the optimal TSC gains.

As a performance index for the comparison of system behavior with different gains, throughput (TP) was calculated [55], here defined as

\[ TP = \frac{1}{n} \sum (1/n) \sum (IDₜ/MTₜ) \]

[bps], where \( n \) is the number of trials and \( m \) is the number of participants. TP is a useful measure for the speed and accuracy performance of movements by the integrated interpretation of the slope and intercept parameters of the regression model. The one-way within-subjects analysis of variance (ANOVA) test was used, and post-hoc analysis was performed using the Tukey’s test.

Table III lists the ANOVA test results, showing a significant effect of the gains for TP (\( p < 0.001 \)), which indicated that the different TSC gains affected the task performance. Fig. 11 shows the TP values of the TSC gains. When \( K₀ = 0.2 \), none of the participants performed better due to the strong constraints of the movements on the guidance path in performing the Fitts’ targeting task. This meant that a greater weighting of the ability of the autonomous controller to control the manipulator hindered the operator’s conscious actions to move the tip out of the guidance path to handle erroneous situations such as failed target detection, target movement during manipulation, and target occlusion by floating particles in the liquid medium. Conversely, a greater weighting on \( K₀ \) means that more manipulation capability is given to the operator; when \( K₀ = 1.0 \), the manipulation is the same as with the direct manipulation mode (no intervention of the controller). In the direct manipulation, the participants easily strayed from the guidance path in performing the steering task and had a lower TP.

Fig. 12 shows the total number of errors for all trials (\( F_{4,12} = 30.4476, p < 0.001 \)). As defined above, the error was counted when the tip failed to track the guidance path (steering law) or in pointing to the target (Fitts’ law). For higher \( K₀ \), the accuracy was lower due to weak constraints on the guidance.

Fig. 10. Relationship between the actual and predicted values (from equations in Table II). The dashed line represents the unit slope for an ideal model.
In this study, the evaluation for $K_h = 0$ could not be given because it was impossible to perform the Fitts’ targeting task due to strong constraints; the operator could only control the manipulator in the guidance path. Although the fully automated system or the intervention of the autonomous function in telemanipulation could enhance microinjection performance by reducing human deficiencies, as the weight of the controller was increased, the operators lost their ability to control the manipulator's motion. Microinjection is a tedious task requiring much skill and practice to manipulate tiny cells with delicate glass tools, which requires cell separation, selection, targeting, tracking and multidimensional manipulation. Therefore, the loss of control ability is problematic for this complicated task. In addition, automatic dexterous manipulation including the rotation of cells has not yet been reported, and there are difficulties in target selection in visual processing. Jang et al. [18] reported that the failed recognition rate in the worst case was 47.8% for the detection of the nucleus and the polar body of mouse embryos. In [19] and [20], approximately 15% of the attached cells (CHO-K1 and endothelial cells, respectively) were missed in the visual targeting process.

IV. CONCLUSIONS

In the context of the single-cell microinjection system, we proposed a TSC to achieve simultaneously high throughput and dexterity. In particular, we provided a quantitative analysis method to determine what level of automation (or direct manipulation) is needed for optimal TSC. Most previous studies on shared control have not addressed how to decide how much autonomy will be integrated in the teleoperation (or vice versa) for the best performance. In this paper, for the evaluation of TSCs with different weightings, the microinjection task was modeled through the Fitts’ and steering laws, which describe the speed/accuracy trade-offs in human movement. The results showed that a 40–60% weighting on the human operator (or the controller) produced better performance for both speed and accuracy of task completion in microinjection. Although the fully automated function could not be measured by the proposed method, the test results suggested some level of automation or human involvement was needed in the development of the robotic biomicromanipulation system from the standpoint of speed and accuracy performance.

In future work, microinjection experiments with biological samples will be performed with clinical evaluations. In addition, different shared control methodologies, such as a
DOF partitioning and a virtual fixture, will be compared for


