Computer Simulation of Low Frequency Vibration in Electrochemical Machining

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ABSTRACT
This research studies how particles transport between low frequency vibrating electrodes during electrochemical machining (ECM). The ANSYS Fluent software was used to study the particle speed while the Star CCM+ software was utilized to study particle interactions during vibration-assisted ECM process. A series of simulations were conducted to calculate the particle average flushing speed. Collided particles either gained momentum or deflected their trajectories to accelerate in the flow of electrolyte. Simulation results showed that the highest average flushing speed of 0.4 m/s was obtained at 40 Hz vibration frequency and 10 µm vibration amplitude. Such higher flushing speed of particles improved machining depth (material removal rate) and produced a sharper machined profile. Experiment results confirmed that the maximum machining depth and minimum taper angle were obtained when vibrating the anodic workpiece at 40 Hz and 10 µm amplitude. Machining depth and ECM material removal rate had a positive correlation with the average flushing speed. A sharper ECM’ed profile was achieved since the taper angle was favorably reduced at high average flushing speed.

Keywords: Low frequency vibration, Simulation, Pulsed current, Electrochemical machining, Particle flushing.

1 INTRODUCTION
Electrochemical machining (ECM), based on the principle of controlled anodic dissolution, has been used to fabricate complex shapes from an electrically conductive component. This is particularly useful to shape engineering alloys that are difficult to be machined such as tool steels, tungsten carbides and superalloys. This ECM process does not induce any thermal or mechanical residual stresses nor subsurface defects below a workpiece, generates no burr, has no tool wear, yet could achieve an excellent surface quality when the parameters are suitable for electrochemical polishing. Since we can eliminate secondary processes for deburring and stress relieving when applying ECM, this process would be a promising technique to machine/polish metallic materials with a decent material removal rate (MRR) can be achieved.

Ion transport mechanism affects the removal rate in ECM. The movement of ions is controlled by three mechanisms: migration, diffusion and convection [1]. A high MRR requires fast ion and by-product movement rates (i.e. by increasing one or all of the migration rate, diffusion rate and convection rate). Researchers have attempted to utilize different techniques to improve these transport rates with different approaches by:

- Applying a high voltage to increase migration rates [2-4]
- Applying pulsed current instead of using direct current [5-8]
- Increasing high electrolyte flow rates to increase diffusion rates [3,9]
- Vibrating either workpiece or electrode at low vibration frequency to increase the convection rates [8,10-14]
- Applying ultrasonic vibration of electrode or via electrolyte [13,15,16].

Convection is the most significant factor that affects ion transport mechanism. The classical Faraday’s law and published literature have shown that a higher voltage/current resulted in high MRR; however, the resulted by-products and excessive heat must be removed, for example, with a costly electrolyte pumping system. An alternative solution is to incorporate pulsed current and vibration into ECM to improve the electrolyte flow while reducing its temperature. Researchers have attempted to vibrate either electrode or workpiece at different vibration frequency

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and amplitude to enhance the process instead of installing an expensive pump for high volume electrolyte flow rate that might deflect a fragile or low compliance workpiece. Although vibration-assisted ECM has been studied experimentally, the mechanism on how low frequency vibration improves MMR is yet to be investigated. The objectives of this paper are to:

i) Simulate the flow of by-product in a horizontally vibrated ECM system.
ii) Study the by-product interaction between electrodes.
iii) Verify the simulation study with experimental data.

2. LITERATURE REVIEW

Many researchers have performed experimental investigation on the effect of vibration on ECM. In an ultrasonic-assisted ECM system, controlled vibration of either the tool electrode or electrolyte to agitate the abrasives suspended in the electrolyte had shown to improve surface finish of the workpiece [16]. A study of the geometry and type of electrode that resulted in a well-polished surface (0.7 µm Ra) was reported and the effect of ultrasonic energy was acknowledged [17]. This energy was also responsible for the removal of debris from the machining zone and creation of optimal hydrodynamic conditions that affected the surface layers [18].

Hybridization ECM with low frequency tool vibration provided a positive and beneficial effect by changing the physical conditions in the inter-electrode gap (IEG). The variation of the gap pressure led to the removal of the sludge products and allowed renewal of the electrolyte in the machining gap. The reciprocal motion between the tool and the workpiece surface enhanced the circulation of the electrolyte through the interface to permit the use of higher current densities in order to improve the quality of the resultant surface and faster removal rate. In traditional vibration-assisted ECM the anode/cathode were vibrated in the vertical direction, thus the resulted by-products, that deposited on either electrode due to gravity, would hinder ion transport mechanism. A horizontal and vibration-assisted ECM was proposed as a remedial solution [6,16,19-21], in which a workpiece vibrated in the horizontal direction instead of the vertical direction to facilitate flushing of the resulted by-products.

For ECM and its variant processes, it is essential to estimate the volume of anode material removal during a specific time. The MRR is a function of current density distribution in the IEG where varying electrical conductivity of the electrolyte exists. Electrolyte velocity and pressure field in the IEG affected the gas bubble formation and temperature distribution, hence degraded the properties of electrolyte properties and current density [22]. As the consequence, any numerical simulation model would include mass, momentum, heat, electric charge and energy balance equations [23,24]. Due to the non-contact nature of the ECM process, it was essential to develop a simulation model to predict the anodic profile. The first electrode shape change simulation model was based on analytical techniques. An ECM tool design was suggested by using the “sine rule” [1]. This rule can be used to obtain an approximate shape but it failed when the tools had sharp discontinuities due to the negligence of stray current effects and the assumption of parallel flux lines. Other authors suggested to use an analytical direct computation for two-dimensional (2D) ECM, represented the workpiece by Fourier’s series and validated their models with experimental work [25], or utilizing finite element method for the 2D computation of MRR in ECM [26,27]. The latter model took the effects of simultaneous changes in the electrolyte flow speed and temperature rise into consideration.

Numerical simulation of three-dimensional (3D) electrode shape changes obtained during the ECM processes based on the “marker” method was also proposed [28]. A complex 3D structure was fabricated by pulsed electrochemical micromachining (PECMM) using ultrashort pulses; The effect of unsteady phenomena in electrical double layer on final anode shape also included in this study [29]. Numerical simulation of the ECM process included the temperature effects was studied in other investigations and the temperature distribution was found to have an influence on the shape of the anode [23,24]. For better accuracy and simplification of tool design, a small yet stable IEG (by reducing the nonuniformity of the electrolyte conductivity) was required. Commercial software package was also used to predict the final anode shape [30].

A review of mass transfer issues and other problems associated with the ECM process was published [31] and the numerical modeling of the ECM process considering the hydrodynamics involved in the process was also studied [32,33]. The final anode shapes resulting after the ECM using a triangular shaped cathode and curved surface were modeled in these studies. Another model was developed to predict the variation of gap in the ECM process using pulsed current with a stationary electrode and rotating workpiece. The model predicted the gap, evolution of anodic profile, and estimation of resulted surface roughness value [34]. Simulation of heat generation during ECM process and its effective dissipation using electrolyte flow was studied; it was found that a hollow cathode and pulse voltages effectively helped to control the heat generation [35].

Published literature did not cover the interaction of discrete particles between electrode gap and the effect of vibration frequency on particle flow in an electrolyte. The following sections describe details of two simulations using Fluent and Star CCM+ software. The simulation results are then compared with experimental and published data.
3. SIMULATION OF PARTICLE FLOW

3.1 Governing Equations

A successful ECM process depends on the efficiency of debris flushing and effectiveness of fresh chemical replenishment to anodic surface. The motion can be modeled using Newton’s second law of motion; the governing equations for translational motion of discrete particle $i$ with mass $m_i$ can be written as:

$$m_i \frac{d\vec{v}_i}{dt} = \sum_j F_{ij} + \sum_k F_{ik}^n + F_{i}^g + \sum c F_{i}^c$$  \hspace{1cm} (1)

Where,

- $\vec{v}_i$: translation velocity of particle $i$ respectively (m/s)
- $F_{ij}$: contact force acting on particle $i$ by particle $j$ (N)
- $F_{ik}^n$: noncontact force acting on particle $i$ by particle $k$ or other sources (N)
- $F_{i}^g$: particle-fluid interaction force on particle $i$ (N)
- $m_i$: mass of particle $i$ (kg)
- $F_{i}^c$: gravitational force (N).

During an ECM process, a typical removal rate of an anode is 45.3 mm$^3$/min which is much less than the electrolyte flow rate of 2.5 L/min (2,500,000 mm$^3$/min) [6]. It is assumed that by-product particle interaction is insignificant, so single particle instead of multiple particles can be used in simulation of debris flushing. Therefore, the contact force and non-contact forces between particles can be neglected, the equation (1) can be simplified to:

$$m_i \frac{d\vec{v}_i}{dt} = F_i^g$$  \hspace{1cm} (2)

Once the forces in equation (2) are known, this equation can be readily solved numerically to determine the particle’s trajectory and velocity.

3.2 Particle-Fluid Interaction Forces

In addition to the buoyancy force, various particle-fluid interaction forces will be generated during the particles interact with surrounding fluid. For example, the translation of particle was driven by drag force and resisted by stagnant fluid. Thus, particle-fluid interaction forces could be considered [36]. Several forces can be implemented in discrete element method in computing fluid dynamics, for examples drag force, pressure gradient force, unsteady force, and lift forces [37,38].

3.2.1. Fluid-Particle Interaction Drag Force

For an isolated sphere particle suspended in a fluid, the Newton’s equation was used to determine the drag resistance force. The particle-fluid drag coefficient $C_d$ is dependent upon Reynolds’s number $Re$, in addition to liquid properties. There are three regions: the Stoke’s Law region, the transition region, and Newton’s law region.

Two methods were used to determine the particle-fluid drag force. The first method based on empirical correlations for either bed expansion [39] or bed pressure drop experiment [40]. The other method based on numerical simulations at microscale, where the technique used the direct numerical simulation [41]. The latter method, although rational, was limited by the current computation capability. Numerical studies had been applied to relatively simple systems. A study that systematically investigated and quantified the differences among these correlations was published [42]: The results revealed that these correlations with similar predictive capability, although their accuracy may differ. In this study, the coefficient was adopted from another study [43], and the fluid-particle interaction drag force are defined as:

$$F_{d} = m \frac{\vec{V}_f - \vec{V}_p}{\tau_r}$$  \hspace{1cm} (3)

$$\tau_r = \frac{\rho_p d_p^2}{18 \mu} C_d Re$$  \hspace{1cm} (4)

$$Re \equiv \frac{\rho_p d_p |\vec{V}_f - \vec{V}_p|}{\mu}$$  \hspace{1cm} (5)

$$C_d = a_1 + a_2 \frac{d_p}{Re} + a_3 \frac{1}{Re}$$  \hspace{1cm} (6)

Where,

- $F_{d}$: drag force (N)
- $\vec{V}_f$: fluid phase velocity (m/s)
3.3. Fluid-Particle Flow Modeling

Simulations using complementary software were employed in this study. The ANSYS Fluent software was used in this study to simulate single particle flow while neglecting particle-particle interaction forces. The Star CCM+ software was used to simulate multiple particles flow which can capture the particle-particle interaction in a fluid-particle flow system.

In the discrete element method of computational fluid dynamics (CFD-DEM), the primary phase (fluid) flow was determined by the CFD on a computational cell scale and the second phase (particles) was obtained by solving Newton’s equations of motion \[44\]. The governing equation for fluid was the Navier-Stoke equation which was the same as the two-fluid method (TFM). The CFD-DEM technique was used in this study because of its superior computational convenience and capability to capture the particle physics, such as, trajectories and velocities. In the CFD-DEM approach, the modeling of particles flow was at individual particle level and the modeling of fluid flow by CFD was at computational cell level. At each step, the DEM would give information, such as locations and velocities of individual particle, for the evaluation of porosity and volumetric fluid drag force in a computational cell. The CFD would then use these data to determine the fluid flow field which then yield the fluid drag forces acting on individual particles. Incorporation of the resulting forces into the DEM will produce information about the motion of individual particles for next time step. At each time step, the fluid-particle interaction forces on individual particles in a computational cell were calculated first, and then the values were added to produce the particle-fluid interaction force at the cell scale, as shown in following equations.

\[
 F_{new} = F_{old} + \alpha (F_{calculated} - F_{old}) \\
 Q_{new} = Q_{old} + \alpha (Q_{calculated} - Q_{old}) \\
 M_{new} = M_{old} + \alpha (M_{calculated} - M_{old})
\]

Where,
\[\alpha \] under-relaxation factor for particles
\[\alpha = 0.9 \text{ for transient flow simulation with unsteady particle tracking} \]
\[\alpha = 0.5 \text{ otherwise} \]
\[F\] force
\[Q\] energy
\[M\] momentum

3.4. Particle Flow Simulations in Interelectrode Gap

This study utilized high strength low alloy (HSLA) steel in experiments to confirm the simulation trends. Since the major composition of HSLA steel is iron (97-99%), during ECM process several possible reactions may occur at the workpiece and electrode. The reaction is the dissolution of iron at anode.

\[Fe \rightarrow Fe^{2+} + 2e^-\]

At the electrode, hydrogen and hydroxyl ions are produced:

\[2H_2O + 2e^- \rightarrow H_2↑ + 2OH^-\]

Thus, the overall reaction of iron in ECM process is:

\[Fe + 2H_2O \rightarrow Fe(OH)_2 + H_2↑\]

The ferrous hydroxide Fe(OH)_2 may further react with water and oxygen to form ferric hydroxide Fe(OH)_3.

\[Fe(OH)_2 + 2H_2O + O_2 \rightarrow 4Fe(OH)_3\]
pulsed ECM. While Star CCM+ was used to simulate how multiple particles would flow and interact in vibration assisted pulsed ECM. The effects of vibration frequency and vibration amplitude on particle average flushing speed are presented in the result section.

3.4.1. Single Particle Simulation

Figure 1 shows the horizontal tubular electrode and workpiece with origin at the tube center. The workpiece surface is on the vertical y-o-z plane and is separated from the tubular electrode at a small interelectrode gap. Consider a by-product particle P between the electrode and workpiece and its free-body diagram. The particle, forming an angle $\varphi$ with the horizontal axis, is subjected to gravity force, buoyance force, and drag force in flowing electrolyte. Due to the axisymmetric electrode, the drag force is in radial direction away from the tube center. The sum of forces in the radial direction is:

$$F_r = F_d - (G - F_b) \sin \varphi$$  \hspace{2cm} (14)

Where,
- $\varphi$: polar angle of the particle
- $F_r$: total force along radial direction (N)
- $F_d$: drag force (N)
- $G$: gravity force (N)
- $F_b$: buoyance force (N)

Since the solid particle is heavier and sinks in electrolyte, the buoyance force of this particle is always smaller than its gravity force. The minimum value of radial direction force equals to $F_d - (G - F_b)$ when $\varphi = \pi/2$, and the maximum value of this force equals $F_d + (G - F_b)$ when $\varphi = 3\pi/2$. The top most location (at $\varphi = \pi/2$) is the most difficult location for solid by-products to be flushed away (location A in Figure 2a). In this study, location A will be taken as the simulation cell zone. The thickness of the electrode is 0.3 mm (Figure 1) and the initial distance between workpiece and electrode is 0.3 mm, thus the simulation zone is a 0.3 mm $\times$ 0.3 mm square box (Figure 2b). Let the new coordinate $xyz$ with origin at the lower left corner of the simulation zone. The electrolyte flow speeds from a pump were set to 2, 3, and 4 m/s and assuming the workpiece vibration follows a sine waveform:

$$x = -A_v \sin(2\pi f t)$$  \hspace{2cm} (15)

Let $x$ be the coordinate of the workpiece, the negative sign means the workpiece movement towards the left and away from the origin.

Where,
- $x$: location of workpiece relative to the fixed origin.
- $A_v$: anode workpiece vibration amplitude (m)
- $f$: vibration frequency (Hz).
In the beginning, the particle center was located at lower left corner (Figure 2b) with particle initial velocity of 0 m/s. The boundary conditions for this study are listed in Table 1. A simulation procedure stopped when the y-coordinate of particle center was larger than 0.3 mm, i.e. when the particle was flushed away from the electrode surface.

| Variables                  | Values          |
|----------------------------|-----------------|
| Vibration frequency $f$ (Hz) | 20, 30, 40      |
| Vibration amplitude $A$ (µm) | 5.0, 7.5, 10.0  |
| Flow speed (m/s)           | 2.0, 3.0, 4.0   |

To investigate the influence of the vibration frequency, vibration amplitude, electrolyte flow speed, particle size and particle locations on average flushing speed, a series of simulations were conducted:

1) The first simulation was used to build a relationship between experimental results and simulation results. Three input levels for each variable were set (Table 1). Thus, total number of simulation run was $3^3 = 27$. 

| Vibration Frequency (Hz) | Vibration Amplitude (µm) |
|--------------------------|--------------------------|
| 10                       | 160.00                   |
| 20                       | 40.00                    |
| 30                       | 17.80                    |
| 40                       | 10.00                    |
| 50                       | 6.40                     |
| 60                       | 4.44                     |
| 70                       | 3.26                     |
| 80                       | 2.50                     |
| 90                       | 1.98                     |
| 100                      | 1.60                     |
| 110                      | 1.32                     |
| 120                      | 1.11                     |
| 130                      | 0.95                     |
| 140                      | 0.82                     |
| 150                      | 0.71                     |
| 175                      | 0.52                     |
| 200                      | 0.40                     |
| 500                      | 0.065                    |
| 1000                     | 0.016                    |
| 5000                     | 0.00064                  |
| 10000                    | 0.00016                  |
3) The second simulation was used to theoretically investigate the maximum flushing speed for the experimental system. The electrolyte flow speed was fixed at 4 m/s, and input current for vibration table was fixed at its maximum value (7 A), the corresponding vibration frequency and vibration amplitude will follow $Af^2 = \text{constant}$ based on the specifications of system and are listed in Table 2.

4) The third simulation was used to investigate the particle size on average flushing speed. The electrolyte flow speed was fixed at 4 m/s, vibration frequency was fixed at 40 Hz, vibration amplitude was fixed at 10 µm and particle sizes range from 2 µm to 10 µm.

The discrete phase formulation used by ANSYS Fluent assumed that (i) the discrete phase (ECM by-product) was sufficiently diluted, and (ii) particle-particle interaction and the effects of the particle volume fraction on the continuous phase were negligible. This implied that the discrete phase must be present at a low volume fraction, typically less than 10-12%.

3.4.2. Multiple Particles Simulation

The Star CCM+ was used to capture the particle-particle interaction in multiple particles fluid-particle flow system. Figure 3 shows the flow domain in $yz$-plane with the length of $wt = 300 \mu m$ and width $La = 20 \mu m$. Four particle injection locations (label “I” and “II” at coordinates $z = \pm 5, y = 50 \mu m$; and label “III” and “IV” at coordinates $z = \pm 5, y = 0 \mu m$) were continuously injecting particles of 2 µm diameter into the electrolyte between the electrode and workpiece.

Due to the limitation of the simulation software, a triangular wave was used as the approximation of the sine wave (Figure 4). During a vibration cycle the workpiece moved at a constant speed of $4Af$ before changing its direction. The by-product particles in ECM were released from the workpiece surface, and their releasing locations changed continuously. To simplify the simulation, a vibration cycle was divided into 20 equal time intervals within which the releasing locations were fixed. When the workpiece moved away from the electrode, particle releasing locations were the same as their initial locations from each time interval; in the reversed direction of workpiece, the particle releasing locations were the ending locations of each time interval.

In this simulation, the vibration was fixed at 40 Hz at 10 µm. The workpiece travel speed was $4Af = 1600 \mu m/s$, and the article releasing rate was $10^5$ particles/s. The complete list of particles releasing locations for one vibration cycle is listed in Table 3.

![Figure 3 Particle injection locations in fluid domain](image)

![Figure 4 (a) Sine wave (b) Approximation of sine wave in Star CCM+](image)
4. EXPERIMENTS

To carry out this investigation, a unique laboratory horizontal vibration-assisted pulsed ECM system was developed (Figure 5). This ECM system can control electrolyte flow, vibration frequency and amplitude, square pulsed DC output and programmable tool feeding. The system (Table 4) included micro workpiece vibration unit, controllable pulsed DC system, programmable feeding system, electrolyte circulation system, and ECM cell with flash guard, etc.

| Simulation Time (ms) | Location I (µm) | Location II (µm) | Location III (µm) | Location IV (µm) |
|----------------------|-----------------|-----------------|-------------------|-----------------|
|                      | \((x_1, y_1, z_1)\) | \((x_2, y_2, z_2)\) | \((x_3, y_3, z_3)\) | \((x_4, y_4, z_4)\) |
| 0 – 1.25             | (1, 50, 5)      | (1, 50, -5)     | (1, 0, 5)         | (1, 0, -5)      |
| 1.25 – 2.50          | (-1, 50, 5)     | (-1, 50, -5)    | (-1, 0, 5)        | (-1, 0, -5)     |
| 2.50 – 3.75          | (-3, 50, 5)     | (-3, 50, -5)    | (-3, 0, 5)        | (-3, 0, -5)     |
| 3.75 – 5.00          | (-5, 50, 5)     | (-5, 50, -5)    | (-5, 0, 5)        | (-5, 0, -5)     |
| 5.00 – 6.25          | (-7, 50, 5)     | (-7, 50, -5)    | (-7, 0, 5)        | (-7, 0, -5)     |
| 6.25 – 7.50          | (-9, 50, 5)     | (-9, 50, -5)    | (-9, 0, 5)        | (-9, 0, -5)     |
| 7.50 – 8.75          | (-11, 50, 5)    | (-11, 50, -5)   | (-11, 0, 5)       | (-11, 0, -5)    |
| 8.75 – 10.00         | (1, 50, 5)      | (1, 50, -5)     | (1, 0, 5)         | (1, 0, -5)      |
| 10.00 – 11.25        | (-1, 50, 5)     | (-1, 50, -5)    | (-1, 0, 5)        | (-1, 0, -5)     |
| 11.25 – 12.50        | (1, 50, 5)      | (1, 50, -5)     | (1, 0, 5)         | (1, 0, -5)      |
| 12.50 – 13.75        | (3, 50, 5)      | (3, 50, -5)     | (3, 0, 5)         | (3, 0, -5)      |
| 13.75 – 15.00        | (5, 50, 5)      | (5, 50, -5)     | (5, 0, 5)         | (5, 0, -5)      |
| 15.00 – 16.25        | (7, 50, 5)      | (7, 50, -5)     | (7, 0, 5)         | (7, 0, -5)      |
| 16.25 – 17.50        | (9, 50, 5)      | (9, 50, -5)     | (9, 0, 5)         | (9, 0, -5)      |
| 17.50 – 18.75        | (11, 50, 5)     | (11, 50, -5)    | (11, 0, 5)        | (11, 0, -5)     |
| 18.75 – 20.00        | (13, 50, 5)     | (13, 50, -5)    | (13, 0, 5)        | (13, 0, -5)     |
| 20.00 – 21.25        | (15, 50, 5)     | (15, 50, -5)    | (15, 0, 5)        | (15, 0, -5)     |
| 21.25 – 22.50        | (17, 50, 5)     | (17, 50, -5)    | (17, 0, 5)        | (17, 0, -5)     |
| 22.50 – 23.75        | (19, 50, 5)     | (19, 50, -5)    | (19, 0, 5)        | (19, 0, -5)     |
| 23.75 – 25.00        | (21, 50, 5)     | (21, 50, -5)    | (21, 0, 5)        | (21, 0, -5)     |
A workpiece plate (#5) was mounted with workpiece surface in vertical direction and an electrode (#6) traveled horizontally into the workpiece with feed rate controlled by a computer-controlled positioner (#7). Workpiece vibration was precisely generated by an electrodynamic shaker (#1). A pulsed power supply (#13) and two high-flow-rate pumps (#11, 16) completed the ECM cell.

In this study, the constant concentration of potassium bromide (KBr, 1mol/L) was selected as electrolyte. Each experimental run was started with filtered electrolyte temperature in the range 21-29°C. Temperature of the electrolyte was measured and recorded by OMEGA HH374 4-channel data logger thermometer. Electrolyte conductivity, measured before each run using the Hannah HI 8733 conductivity meter, was in the range of 111-121 mS/cm. During experimental, fresh electrolyte was pumped from fresh electrolyte container (#16) and flowed inside the cathodic electrode (#4). After machining, used electrolyte in the ECM cell (#5) was pumped out and stored in used electrolyte container (#8). Although the metallic by-products in used electrolyte could be filtered using centrifugal method in small quality, the settling method was used for a large quantity of used electrolyte. In the latter method the used electrolyte, mixed with metal debris and by-products, was stored overnight so that the heavy metallic and salt-byproduct would be settled at the contained bottom. The clear electrolyte at the top was pumped back to the fresh electrolyte container (#16) using a Longer WT600-2J peristaltic pump (#15) and reused for subsequent experiment if its conductivity was still comparable with that of the fresh electrolyte.

The EVERLAST 255 EXT DC power supply provides either DC or pulsed DC of 3-150 A up to 500 Hz pulsed current frequency. Since a high peak current may generate sparks that damage the workpiece and electrode, the constant peak current was set at 26 A, current frequency at 500 Hz and 50% duty cycle.

Stainless steel tubes (Ø9.5 mm OD, 0.3 mm thick, #4) were selected as cathodic electrode. About 13 mm ends were commercially coated with Teflon to form a nonconductive layer of 0.02 mm thick on both outside and inside diameters. The coated ends were carefully sanded off using 600-grit abrasive paper to make the cathodic electrode tube conductive.

The Labworks ET-132-2 electrodynamic shaker was powered by the Labworks PA-151 linear power amplifier that in turn controlled by an Agilent 33250A waveform generator. The computer-controlled positioner included two Velmex motorized frames, a rotary plate and a VXM-3 controller system. The system has a load capacity of 15.9 kg horizontally and 4.5 kg vertically with straight-line accuracy of 0.076 mm/25 cm, feed rate range 2.5-5000 µm/s, and repeatability of 0.0025 mm.

The TENMA 72-6202 multimeter was used to measure the conductivity between the cathodic electrode and the anodic workpiece. The Velmex computer-controlled positioner was used to set the initial inter-electrode gap. At first, the multimeter was used to find the location of the cathodic electrode where in contact with the anodic workpiece since the resistance between the cathodic electrode and the anodic workpiece dropped to 0 (IEG = 0). After that, the cathodic electrode moved back 0.3mm and controlled by the Velmex positioner. In this investigation, a constant feed rate was set to 15µm/s and the cathodic electrode traveled a distance of 2.5 mm. All experimental conditions are summarized in Table 5.

After ECM’ed, all samples were rinsed and cleaned with water in an ultrasonic bath for one minute and then dried with compressed air. All samples were positioned in a grass beaker in the bath with ECM’ed holes facing down to facilitate removal of residual particles from the hole. The Alicona Infinite Focus 3D profiler was used to analyze the ECM’ed machined depth, surface finish and wall taper angle.

The settled by-products were collected and washed with distilled water 10 times to remove the dissolved potassium bromide. Two samples of by-products were prepared for SEM and EDX. The sample for SEM was one drop of by-products solution placed on the titanium plate and dried in air. Thin Au-Pd layer was sputtered coated on the surface of the titanium and specimens. The Tescan Vega 3 SEM was used to capture the surface image of the by-products. The ECM sediment was washed, dried, and deposited on a carbon tape. The Zeiss EDX was used to analyze the by-products compositions.
Table 5 Summary of experimental conditions

| Variables                        | Values                  |
|----------------------------------|-------------------------|
| Current frequency (Hz)           | 500                     |
| Electrode feed rate (µm/s)       | 15                      |
| Electrode travel distance (mm)   | 2.5                     |
| Electrolyte concentration (mol/L)| 1                       |
| Electrolyte conductive (mS/cm)   | 111-121                 |
| Electrolyte flow rate (L/min)    | 2.5                     |
| Starting IEG (mm)                | 0.3                     |
| Peak current (A)                 | 26                      |
| Vibration amplitude (µm)         | 0, 2.50, 4.44, 5.00, 7.50, 10.00 |
| Vibration frequency (Hz)         | 0, 20, 30, 40, 60, 80   |

5 RESULTS AND DISCUSSIONS
Simulation results from ANSYS Fluent for flushing speed, the effect of vibration frequency, and electrolyte flow rate are presented first. The particle interaction results from STAR CCM+ simulation are then followed. Experimental trends are finally presented and compared with the simulation data.

5.1. Average Flushing Speeds
Recall that the simulation cell is 0.3 mm × 0.3 mm (Figure 6a), and a simulation will be terminated if the y-coordinate of a particle center is larger than 0.3 mm, i.e., a particle is effectively flushed and traveled outside of the interelectrode gap. Typical particle trajectory along y-direction as a function of time is shown in Figure 6. The average flushing speed, $V_{\text{ave}}$, is defined as the secant slope of particle path:

\[
V_{\text{ave}} = \frac{\text{particle travel distance}}{\text{travel time}} = \frac{300 \, \mu \text{m}}{756 \, \mu \text{s}} = 0.396 \, \text{m/s}
\]  

The effect of vibration frequency on average flushing speed when the driving current for vibration table was set at $I_{\text{vibration}} = 7A$ is shown in Figure 7. The average flushing speed decreases with increasing vibration frequency. Neglecting the effect of the workpiece vibration on the particle movement along the cathodic electrode x-direction, the distance between workpiece and the particle center is given in Equation (17) and illustrated in Figure 8.

\[
d_g = -A_v \sin(2\pi ft) + r_p
\]  

Where,
- $d_g$ : Gap between particle and workpiece surface ($\geq r_p$) (µm)
- $A_v$ : Vibration amplitude (µm)
- $f$ : Vibration frequency (Hz)
- $r_p$ : Particle diameter (µm)
Figure 7 Effect of vibration frequency on average flushing speed. (Simulation with ANALYS Fluent at $I_{vibration} = 7A$)

As shown in Figure 7, $I_{vibration}$ is fixed at 7 A, the average flushing speed decreases from 0.5736 m/s (at $f = 10$ Hz, $A_v = 160 \mu m$) to 0.3231 m/s (at $f = 10,000$ Hz, $A_v = 0.16 \mu m$) when vibration frequency increases from 10 Hz to 10,000 Hz. For fixed vibration current or constant vibration power, the vibration amplitude and vibration frequency follow the relationship $A_v f^2 = constant B$, and Equation (17) becomes:

$$d_g = -B \frac{\sin(2\pi ft)}{f^2} + r_p$$  \hspace{1cm} (18)

Equation (18) suggests that the distance between particle center and workpiece $d_g$ decreases drastically with increasing vibration frequency $f$. Therefore, for fixed current on vibration generator, a high vibration frequency would lead to a smaller gap, lower drag force and lower average flushing speed, as shown in Figure 7.

The relationship between particle size and average flushing speed is shown in Figure 9. As the particle radius increases from 1 µm to 3 µm, the average flushing speed increases from 0.29 to 2.77 m/s. However, continue increase particle size to 5 µm, the average flushing speed changes slightly, from 2.77 to 3.08 m/s. This is because the gap $d_g$...
in Equation (18) increases with an increasing of particle radius. Larger gap results in higher drag force and higher average flushing speed.

![Figure 9 Relationship between average flushing speed and particle radius. (Simulation with ANASYS Fluent at \( f = 40 \text{ Hz} \) and \( A_v = 10 \mu m \) and \( V_e = 4 \text{ m/s} \)](image)

This simulation results were confirmed with experiment results conducted in another study [12]. For a preset machining parameter combinations (4 µm, 6 µm and 8 µm vibration amplitude, 160 µm tungsten cathodic electrode with 5 µm insulate layer, 321 stainless steel workpiece with 0.5 mm thickness, 5 wt% NaNO\(_3\) + 0.8 wt% EDTA-Na\(_2\) electrolyte, 6 V voltage with 50% duty cycle, 2 kHz pulsed voltage and 15.6 µm IEG), after obtaining the maximum MRR at a 50 Hz, the MRR decreased with increasing of vibration frequency, as shown in Figure 10.

![Figure 10 Effect of vibration frequency on the MRR, reprinted from [12].](image)

5.2. Effect of Vibration on Average Flushing Speed and Part Quality

The effects of vibration frequency and amplitude on average flushing speed, depth of material removal, and taper angle are presented in this section. Figure 11a,b show how the flushing speed and machined depth when varying the vibration parameters. At every vibration amplitude, the average flushing speed increases as the vibration frequency increases. The maximum average flushing speed, 0.396 m/s, is obtained at 40 Hz vibration frequency and 10 µm vibration amplitude. For a specific vibration amplitude \( A_v \), when the vibration frequency increasing, the gap between the anodic electrode surface and the particle center is increased, hence the drag force exert on the particle is increased which result in an increasing average particle flushing speed.

At every vibration amplitude (5, 7.5, and 10 µm), a higher vibration frequency leads to faster average flushing speed and deeper machining depth.

- When vibration frequency increases from 20 to 40 Hz, the average flushing speeds increase by 5.4% (from 0.3423 to 0.3608 m/s), 7.8% (from 0.3517 to 0.3790 m/s) and 9.9% (from 0.3610 to 0.3968 m/s), at respective vibration amplitude.
- Improving of flushing speed changes machining depths by 2.9 % (from 1282 to 1319 µm), 9.9% (from 1308 to 1437 µm) and 18.6% (from 1336 to 1584 µm), respectively.
Vibration-assisted ECM increases the machined depth of a hole while reducing the hole taper angle, therefore, enhancing the product quality. The effects of vibration frequency on average flushing speed and taper angle are illustrated in Figure 12a,b.

- At every vibration frequency (20, 30, and 40 Hz), a higher vibration amplitude leads to faster flushing speed and smaller taper angle—therefore, straighter and sharper hole profile. When vibration amplitude increases from 5.0 to 10.0 µm, the taper angles decreases by 8.7% (from 31.92 to 29.15°), 12.8% (from 29.32 to 25.56°) and 43.4% (from 29.28 to 16.57°), at respective vibration frequency.

- At every vibration amplitude (5, 7.5, and 10 µm), a higher vibration frequency leads to a smaller taper angle. When vibration frequency increases from 20 to 40 Hz, the taper angles decrease by 3.4% (from 31.92 to 29.28°), 27.3% (from 29.81 to 21.67°) and 43.2% (from 29.15 to 16.57°) at corresponding vibration amplitude.
The simulation results and experimental data trends presented above agree with other studies. Simulation and experimental data suggest a negative correlation between average flushing speed and taper angle. The similar trend was also reported on hole shape after ECM (using brass tube cathodic electrode with $\phi8$ mm outer diameter and $\phi3.5$ mm inner diameter, 20 g/L NaCl electrolyte, 6 L/min electrolyte flow rate, 1 mm/min cathodic electrode feed rate, 18 V applied voltage and 50 Hz vibration frequency [11]); the conicity (slope of a cone) decreased from 3.95 to 3.25% when vibration amplitude increased from 20 to 100 µm. This trend can also be explained qualitatively:

i. Vibration of workpiece leads to a higher flushing speed that breaks up agglomerated groups of by-products into smaller chunks of particles.

ii. The smaller chunks forming between two electrodes then travel up the wall of hole and out side of the interelectrode zone. The smaller coalesced particles would erode the wall less and result in smaller taper angle and sharper hole profile.

Low frequency vibration of workpiece increases the flushing speed, enhances ion transport rate and leads to more effective machining rate. Similar results were also reported in other experimental studies in which an ECM system utilized $\phi160$ µm tungsten cathodic electrode with 5 µm insulate layer, 321 stainless steel anodic workpiece with 0.5 mm thickness, 5 wt% NaNO$_3$ + 0.8 wt% EDTA-Na$_2$ electrolyte, 6 V voltage with 50% duty cycle, 2 KHz pulsed voltage and 15.6 µm IEG, 3-14 µm vibration amplitude, and 50-200 Hz vibration frequency [12]. That study reported that:

i. At any vibration amplitude (4, 6, and 8 µm), the MRR increased when increasing vibration frequency from 0 Hz to 50 Hz. However, the MRR dropped when increasing frequency further to 200 Hz.

ii. At both vibration frequencies (50 Hz and 100 Hz), the MRR increased with increasing vibration amplitude in the range 0-8 µm. The maximum MRR was obtained when vibrating at 8 µm amplitude for any vibration frequency.

5.3. Effect of Electrolyte Flow Rate on Average Flushing Speed
The effect of electrolyte flow rates on particle average flushing speeds is illustrated with simulation data (Figure 13a,b).

- At 20 Hz and 5 µm vibration, when the electrolyte flow rate increases from 2 to 4 m/s, the average flushing speed increases from 0.152 to 0.342 m/s.
- At 40 Hz and 10 µm vibration, the average flushing speed increases from 0.198 to 0.396 m/s.
- For each combination of vibration frequency and vibration amplitude, the average flushing speed seems to be proportional to the electrolyte flow rate.

For a single sphere particle in a fluid, Newton’s equation is used to determine the drag resistance force. The particle-fluid drag coefficient, $C_d$, is dependent upon Reynold’s number, $Re$, in addition to liquid properties. There are three regions: the Stoke’s Law region, the transition region and Newton’s law region. By adopting the coefficients from another study [43], the fluid-particle interaction drag force can be modeled as:

$$\bar{F}_f = \bar{F}_d + \bar{F}_p = m \left( \frac{\bar{V}_f - \bar{V}_e}{\tau_f} - \bar{V}_i \bar{\rho}_e \bar{g} \right)$$

$$\tau_r = \frac{\rho_p d_p^2}{18 \mu} C_d Re$$

"Figure 12b Effect of vibration frequency on average flushing speed (left vertical scale) and taper angle (right vertical scale)"
\[ R_e \equiv \frac{\rho_F \mu d_p |\vec{V}_p - \vec{V}_F|}{\mu} \]  
\[ C_d = a_1 + \frac{a_2}{R_e} + \frac{a_3}{R_e^2} \]

Where,
- \( F_f \): Particle-fluid interaction force on particle (N)
- \( F_d \): Drag force (N)
- \( F_b \): Buoyance force (N)
- \( m \): Particle mass (g)
- \( V_f \): Fluid phase flow velocity (m/s)
- \( V_p \): Particle flow velocity (m/s)
- \( V_i \): Translation velocity of particle \( i \) respectively (m/s)
- \( \rho_f \): Fluid density (g/mm\(^3\))
- \( \rho_p \): Particle density (g/mm\(^3\))
- \( \tau_r \): Particle relaxation time
- \( d_p \): Particle diameter (m)
- \( \mu \): Viscosity of the fluid (Pa\(\cdot\)s)
- \( C_d \): Drag coefficient
- \( R_e \): Reynold’s number

For an increasing fluid velocity \( V_f \), the particle-fluid interaction force on particle \( F_f \) is increased, which leads to an increasing particle average flushing speed \( V_{ave} \).

**Figure 13a** Effect of electrolyte flow speed on particle average flushing speed at every vibration frequency.

**Figure 13b** Effect of electrolyte flow speed on particle average flushing speed at every vibration amplitude.
5.4 Multiple Particle Interactions

This section presents the simulation results of possible interaction among ECM particles within an interelectrode gap. Figure 14 illustrates the initial conditions of the simulation. The blue color bar on the left demonstrates the “Particle Residence Time;” the segment (a) shows the anodic workpiece in the yz-plane and the cathodic electrode (not shown) is in the yz-plane toward the positive x-direction. The four green points, I, II, III and IV, represent the particle injection locations. The enlarged view of zone A is shown in segment (b), while segment (c) is the side view of segment (b).

To visualize and follow the motion of a particle at different time and location, define the particle with nomenclature $P_{X,i}(t_j)$: where $X = I, II, III, IV$ is the particle injection location, $i = 1, 2, 3, \ldots, n$ is the particle injection order, and $j = 1, 2, 3, \ldots, n$ is injection time. For example, the particle $P_{IV,1}(t_1)$ represents the 1st particle released at location IV and at time $t_1$. For simplification, only the right side (locations II & IV) is presented since there is no interaction of particles releasing from II and IV locations with those released at the I and III locations.

– At time $t_1 = 1 \mu$s, the particle $P_{IV,1}(t_1)$ is just released from the anodic plate and it is still in contact with the surface (Figure 15).
The timely release of other particles includes:

- The particle $P_{IV,2}$ is released into simulation domain at time $t_2 = 71 \mu s$, represented as $P_{IV,2}(t_2)$
- Particle $P_{IV,3}$ at time $t_3 = 81 \mu s$
- Particle $P_{II,4}$ at time $t_4 = 91 \mu s$
- Particle $P_{II,5}$ at time $t_5 = 141 \mu s$
- Particle $P_{II,6}$ at time $t_6 = 151 \mu s$
- Particle $P_{II,7}$ at time $t_7 = 211 \mu s$

Recall from Figure 2 that the simulation zone is at the top of a workpiece with $300 \mu m$ zone, and the electrolyte flow is upward along the positive Y axis. Thus, the motion of all particles will be in the upward direction of this horizontal ECM setup.

- At time $t_7 = 211 \mu s$, the locations of these particles are illustrated in Figure 16.

![Figure 16](image) The Star CCM+ simulation results at $t_7 = 211 \mu s$.

As simulation continues, there are collisions occur between the particles. Figure 17 and the closed-up Figure 18 illustrate the interaction among particles $P_{IV,2}$, $P_{IV,3}$ and $P_{II,7}$.

- At time $t_8 = 272 \mu s$, the particle $P_{II,7}$ impacts the particle $P_{IV,3}$ (marked with “*”). After collision, this particle $P_{IV,3}$ obtains an acceleration along the y-direction and its flow speed along the y-direction is increased. The particle $P_{II,7}$ obtains an additional acceleration along the x-direction and it moves toward the cathodic electrode (not shown, but on the yz-plane in the positive x-direction), while it continues to move along y-direction due to the drag effect of the fluid.

- At time $t_9 = 280 \mu s$, the particle $P_{IV,3}$ impacts the particle $P_{IV,2}$. After collision, the particles $P_{IV,3}$ and $P_{IV,2}$ will combined and move together. The speed of particle $P_{II,7}$ along the y-direction is increasing due a higher electrolyte flow speed away from the anodic surface.

- At time $t_{10} = 287 \mu s$, after colliding among particles $P_{IV,2}$, $P_{IV,3}$ and $P_{II,7}$, the particle $P_{II,7}$ passes the particle $P_{IV,3}$.

- At time $t_{11} = 291 \mu s$, the particle $P_{II,7}$ passes the particle $P_{IV,2}$, and continue passes the particle $P_{II,6}$.

- At time $t_{12} = 322 \mu s$, the particle $P_{II,7}$ impacts the particle $P_{II,5}$ and continue moves along the y-direction.

- At time $t_{13} = 353 \mu s$, the particle $P_{II,7}$ has already passed the particle $P_{IV,1}$. 
Figure 17 Illustration of particle collisions within zone B from $t_{8} = 272 \mu s$ to $t_{13} = 353 \mu s$.

Figure 18 Enlarged view of Figure 17. A “*” is marked on particle $P_{IV,3}$ for identification purpose.

Figure 19 illustrates the motion of particles $P_{X,1-6}$ from $t_{14} = 512 \mu s$ to $t_{16} = 940 \mu s$.

- At time $t_{14} = 512 \mu s$, the $y$ coordinate of the particle $P_{II,7}$ is larger than 300 $\mu m$, which means that the particle is flushed away from the interested simulation domain. The particle $P_{IV,2}$ obtains an acceleration along the $y$-direction due to a collision and then its speed is larger than that of the particle $P_{IV,3}$. Hence, these two particles are separated and move independently.

- At time $t_{19} = 940 \mu s$, the $y$-coordinate of the particle $P_{IV,1}$ is larger than 300 $\mu m$, as shown in Figure 19 and 20. There is no interaction of particles releasing from II and IV locations with these from I and III locations. Thus, it is reasonable to use the right side of the workpiece to analysis the multiple particles flushing process.

The average flushing speed of the particle $P_{IV,1}$ is:

$$V_{ave} = \frac{\text{total travel distance}}{\text{travel time}} = \frac{300 \mu m}{940 \mu s} = 0.319 \text{ m/s}$$  \hspace{1cm} (23)

The average flushing speed, calculated from Star CCM+ (0.319 m/s showing above), agrees with the result calculated from the ANASYS Fluent (0.396 m/s) shown in equation (16).
5. CONCLUSIONS AND RECOMMENDATIONS

This research used both the ANSYS Fluent and Star CCM+ software to simulate particle transport process within interelectrode gap in the low frequency vibration-assisted electrochemical machining (ECM) process. A series of simulations were conducted to calculate the particle average flushing speed. The study aimed to find relevant process parameters to increase particle flushing speed, therefore, material removal rate and part quality after ECM. This study showed that:

1) Simulations of the motion of a single particle using computational fluid dynamic are performed to track the particle motion and speed between two electrodes. Flushing of the by-products is characterized by...
calculating of particle flushing speed when it exits inter-electrode gap. The maximum flushing speed of ~0.4 m/s was calculated for workpiece vibration frequency of 40 Hz at 10 µm vibration amplitude.

2) Since the particle generating rate is significant smaller than the electrolyte flow rate, it is reasonable to use a single particle for simulation instead of involving multiple particles in the simulation.

3) The simulation results indicate the average flushing speed in ECM would increase with: workpiece vibration frequency, vibration amplitude, electrolyte flow rate, and particle size.

4) The ECM experimental results show that the high flushing speed of by-products would enhance the process since it produces a deeper hole (increasing material removal rate) while forming a sharper hole profiles (increasing product quality).

5) The 2 µm spherical particles were used in this study, further study should consider the interactions between different shapes and different size of by-product particles.

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