Plasmofluidic Microlenses for Label-Free Optical Sorting of Exosomes

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Optical chromatography is a powerful optofluidic technique enabling label-free fractionation of microscopic bioparticles from heterogenous mixtures. However, sophisticated instrumentation requirements for precise alignment of optical scattering and fluidic drag forces is a fundamental shortcoming of this technique. Here, we introduce a subwavelength thick (<200 nm) Optofluidic PlasmonIC (OPTIC) microlens that effortlessly achieves objective-free focusing and self-alignment of opposing optical scattering and fluidic drag forces for selective separation of exosome size bioparticles.

Our optofluidic microlens provides a self-collimating mechanism for particle trajectories with a spatial dispersion that is inherently minimized by the optical gradient and radial fluidic drag forces working together to align the particles along the optical axis. We demonstrate that this facile platform facilitates complete separation of small size bioparticles (i.e., exosomes) from a heterogenous mixture through negative depletion and provides a robust selective separation capability for same size nanoparticles based on their differences in chemical composition. Unlike existing optical chromatography techniques that require complicated instrumentation (lasers, objectives and precise alignment stages), our OPTIC microlenses with a foot-print of 4 μm × 4 μm open up the possibility of multiplexed and high-throughput sorting of nanoparticles on a chip using low-cost broadband light sources.

Optical chromatography is a powerful fractionation technique that has recently gained significant attention for label-free separation and analysis of microscopic bioparticles (cells, bacteria, etc.)4–6. This technique relies on a mildly focused Gaussian laser beam along a microfluidic channel to create opposing optical scattering and fluidic drag forces5,6. One can harness the varying strength and balance of these forces on bioparticle size, composition and morphology for selective sorting. Initially demonstrated for size-based elution of polystyrene beads, optical chromatography has been successfully applied to fractionation of blood components such as human erithrocytes, monocytes, granulocytes, and lymphocytes2,5. This technique particularly excels in distinguishing bioparticles with subtle differences4–7. Its highly precise separation capability has been recently demonstrated using micron size bioparticles with diameters that differ by less than 70 nm8. Most remarkably, separation of two closely related genetic relatives, Bacillus anthracis and Bacillus thuringiensis4, and cells with single gene modifications3 has been achieved using small differences in their chemical makeup (refractive index). In essence, optical chromatography offers unique capabilities as a modern separation technique, especially when combined with multi-stage sequential fractionation and microfluidic network-based purification approaches9,10. However, several critical limitations must be overcome for its wide adaptation: (1) To create strong optical scattering forces along the microfluidic channels, high cost laser sources are needed6. (2) Laser beam must be precisely aligned along the fluidic channel with a well-controlled beam waist profile, requiring a complicated optical alignment procedure that employs multiple multi-axis positioners5. (3) Scaling of existing optical chromatography techniques for multiplexed and high throughput operation is not practical, since each channel requires separate alignment sensitive and high-cost laser sources and objectives10.

In this report, we introduce Optofluidic PlasmonIC (OPTIC) microlenses to overcome the shortcomings of existing optical chromatography techniques by eliminating the need for sophisticated instrumentation and precise alignment requirements. Our sub-wavelength thick (~200 nm thick) OPTIC microlenses offer objective-free focusing and self-alignment of optical and fluidic drag forces and present a facile platform for selective separation.
of exosome size bioparticles. By allowing direct coupling of collimated broadband light to realize strong optical scattering forces at a focal point, extremely small footprint (4 μm × 4 μm) OPtIC microlenses open the door for drastically multiplexed optical chromatography and high-throughput sample processing capability. In addition to parallel operation, laterally integrated OPtIC microlenses on a planar chip permit serial microfluidic schemes9,10 to be readily implemented for multi-stage sequential separation and purification using broadband light sources and conventional planar microfluidic approaches. In the following, we show that our optofluidic sorting scheme based on OPtIC microlenses enables selective separation of exosome size bioparticles (<200 nm in diameter). Furthermore, we demonstrate that our platform offers readily tunable, highly reliable and selective separation of nano-bioparticles by adjusting the light intensity (i.e. radiation pressure) and/or the fluid flow rate (i.e. opposing drag force) based on size and minor difference in chemical makeup (refractive index).

**Results**

Optofluidic-plasmonic (OPtIC) microlenses. As shown in recent studies, periodic and quasi-periodic arrays of plasmonic nanoapertures can be used as micro-convex lenses focusing incident plane waves to dimensions comparable to optical wavelengths11–13. In addition to significantly-small fingerprints, this focusing capability can be exploited to realize large optical scattering forces using collimated broadband light sources14,15. In particular, finite size nanohole array (NHA) structures provide a broadband focusing capability that is mainly controlled by the overall size of the array and tolerant of substructural variations11. The distinct nature of light focusing mechanism in NHA microlenses opens the door for nanofluidic integration through alterations in array design without degrading their focusing characteristics. A suspended plasmofluidic (OPtIC) microlens based on this principle is shown in Fig. 1. Here, an 9 × 9 array of nanoholes with a diameter of d = 150 nm and a periodicity of a₀ = 380 nm is considered (Fig. 1a). NHA patch with a foot print of 4 μm × 4 μm is defined through a 120 nm thick gold film, 10 nm thick accompanying titanium (Ti) adhesion layer and a suspended silicon nitride (Si₃N₄) membrane (thickness of 100 nm) for microfluidic access from both side of it16. The periodic nanohole lattice allows enhanced light transmission through extraordinary light transmission effect (EOT)17–20. The center nanohole is enlarged to facilitate nanofluidic flow. In Fig. 1a, the enlarged center hole diameter is d_c = 500 nm. The overall device is a hybrid nanoplasmonic and nanofluidic system in which fluidic flow through the enlarged center aperture is achieved by pumping a solution from an inlet port on the top side and extracting it through an

![Figure 1](https://example.com/image1.png)

**Figure 1.** (a) A 4 μm × 4 μm OPtIC microlens consisting of a patch array of small circular nanoholes and an enlarged center aperture for nanofluidic integration. (b) Fluidic flow profile through the OPtIC microlens with a fluidic flow rate of 1.3 μm/s at the focal point. (c) Separation of two different size particle is illustrated at the focal point region. Fluidic drag (F_d), optical scattering (F_s) and thermo-plasmonic convection (F_tp) forces are inherently aligned against each other along the optical axis by the OPtIC microlens.
outlet port on the bottom side21,22. The inlet and outlet ports are located away from the plasmofluidic microlens to provide a clear path for the optical beam11. Following Hagen-Poiseille's law \( Q = \Delta P / R_H \) \( [m^3 \cdot s^{-1}] \), the pressure-driven flow through a circular opening of length \( h \) can be understood using hydraulic resistance \( R_H \), which is inversely proportional to the fourth power of the opening radius \( r \): \( R_H \approx 8 \mu \eta / \pi r^4 \) \( [Pa \cdot s^{-1}] \), where \( \mu = 8.9 \times 10^{-4} \) \( Pa \cdot s^{-1} \) is the dynamic viscosity of water and \( \eta = \rho g + r + \eta \) \( 205 \) \( mm \) is the cylindrical conduit thickness. For an enlarged circular center aperture of \( d_c = 500 \) \( nm \) the hydraulic resistance shows nearly an order of magnitude smaller than that of the smaller nanoholes \( d = 150 \) \( nm \) around it. The flatness hydraulic resistance path through the enlarged center aperture \( d_c = 500 \) \( nm \) leads to focusing of the convective fluid flow along optical axis (OA), as demonstrated in our finite-element method (FEM) microfluidic calculations (Fig. 1b). Here, the flow through the centrally located enlarged aperture serves two purposes: (1) it brings nanoparticles to the focal point of the OPTIC microlens, and (2) it forces nanoparticles to follow a trajectory that goes through the OA, as illustrated in Fig. 1c. One can achieve optofluidic alignment by introducing a collimated light beam that is perpendicularly incident from the bottom of the OPTIC microlens (Fig. 1c). The beam focused by the OPTIC microlens, hence optical scattering force \( F_s \), is spontaneously aligned against the fluidic drag force \( F_D \) along the OA. Electromagnetic heating of the plasmofluidic microlens establishes a temperature gradient and heat induced convection current. The resulting thermo-plasmonic drag force \( F_D \) contributes to the optical scattering processes in balancing the fluidic drag. As illustrated in Fig. 1c., in addition to providing a scattering force \( F_s \) along the OA, light focused through the OPTIC microlens introduces an optical gradient force \( F_O \) that radially pushes particles towards the OA. Combined with the radial drag forces \( F_D \) at the focal point region, optical gradient force provides a robust mechanism for precise alignment of particles along the OA, as illustrated in Fig. 1c. For sufficiently large optical pressures that overcome the fluidic drag force \( F_D > F_O \), particles are propelled against the fluidic flow along the OA. Hence, larger and higher refractive index particles (Fig. 1c, green color) experiencing larger optical scattering forces are kept at a distance away from the planar surface, whereas the smaller diameter and lower refractive index particles (Fig. 1c, yellow colored) are allowed to pass through the enlarged center aperture, resulting in complete separation of small size bioparticles (i.e., exosomes) from a mixture of larger size particles through negative depletion. Here, the OPTIC microlens serves as a far-field screen that prevents clogging of the center nanofluidic channel by keeping larger bioparticles away from the nanosurface aperture and high flow rate regions close to the center aperture.

**Light focusing with plasmofluidic microlens.** A critically important device parameter is the nanofluidic channel (center aperture) diameter, which controls the size exclusion range in addition to the focusing behavior of the OPTIC microlenses13. The enlarged center aperture \( d_c \) must be incorporated without deteriorating the focusing characteristics and the desired size selectivity. In Fig. 2a, intensity profiles of OPTIC microlenses with varying center aperture diameters are shown for an incident light beam at \( \lambda = 633 \) \( nm \). The intensity profiles of a NHA without a center aperture \( d_c = 0 \) (a) a uniform NHA \( d_c = 150 \) \( nm \) and a NHA with an enlarged center aperture \( d_c = 500 \) \( nm \) reveals that the enlarged center aperture has negligible effect on the focusing behavior. Even for significantly larger center aperture dimensions \( d_c = 800 \) \( nm \), OPTIC microlens manages to focus light to a tight spot, albeit with a lower efficiency (degraded focusing characteristics). The checkerboard-like intensity profile close to the microlens surface (see Supplementary, Fig. S1) is due to the diffractive self-imaging of smaller size nanoholes around the center aperture, an effect known as plasmonic Talbot effect13,24. The enhanced light intensity regions around the larger diameter center openings \( d_c = 500 \) \( nm \) and \( 800 \) \( nm \) is associated to the diffractive transmission of light through the enlarged aperture. As shown in Fig. 2b, for large enough openings \( d_c \) \( \geq 500 \) \( nm \), the light intensities in these diffractive transmission regions are comparable to or higher than those at the focal point. However, in contrast to the focal point, fluidic drag forces \( F_D \) \( \propto v \) are much stronger than the optical scattering forces \( F_s \) \( \propto v \) in these diffractive transmission regions. As a result, orders of magnitude enhanced fluidic flow velocities close to the center aperture as the fluidic flow squeeze through the narrow center aperture \( d_c = 500 \) \( nm \) diameter (Fig. 1b). Hence, small size and lower refractive index particles that are filtered through the focal point region can follow the fluidic flow lines to the other side of the OPTIC microlens without hinderance in the diffractive transmission regions close to center aperture. Therefore, in the following, we optimized our OPTIC microlens design for the focal point where comparable strong optical scattering, thermo-plasmonic convection and fluidic drag forces can be readily created for selective sorting purposes.

Enlarged center aperture minimally affects the focal distance \( f_o \) as shown in Fig. 2b. The focal distance \( f_o \) is \( 5.32 \) \( \mu m \) for \( d_c = 0 \), \( 150 \) \( nm \), and \( 500 \) \( nm \), whereas it only slightly increases to \( 5.56 \) \( \mu m \) for \( d_c = 800 \) \( nm \). Similarly, a wider focal spot size at \( z = f_o \) is observed for microlenses with larger center apertures (Fig. 2c). For microlenses with \( d_c = 0 \) and \( 150 \) \( nm \), spot sizes are almost identical with a full-width at half maximum (FWHM) spread of \( w = 1.12 \) \( \mu m \). For \( d_c \geq 500 \) \( nm \), \( w \) increases with increasing center aperture opening; for \( d_c = 500 \) \( nm \) and \( 800 \) \( nm \), we find that the spot sizes are \( w = 1.24 \) \( \mu m \) and \( 1.80 \) \( \mu m \), respectively. Focusing characteristics of OPTIC microlenses are mainly controlled by the smaller diameter nanohole array created around the center enlarged aperture. Transmission spectra obtained using a field monitor is shown in Fig. 2d. EOT effect occurs at wavelengths that satisfy the Bragg condition: \( G = iG_i + jG_j \), where \( (i, j) \) are the integer values for the corresponding grating order13,19. In Fig. 2d, the gratings coupled (1,0) resonance transmission peak is observed at \( 650 \) \( nm \) (FWHM \( \sim 55 \) \( nm \)), confirming that light focusing is a result of interference of in-phase photons emerging from the periodic small size nanohole array21. For center apertures with diameters up to \( d_c = 500 \) \( nm \) (Fig. 2d, yellow curve), transmission spectra show minimal deviation from a uniform NHA microlenses without a center aperture \( d_c = 0 \)13, albeit with some broadband background transmission. On the other hand, for significantly larger center openings \( d_c = 800 \) \( nm \), non-resonant light transmission through the center aperture increases, leading to an increased background signal and broadening of the resonant transmission peak (Fig. 2d, violet curve). Based on the focusing behavior, attainable fluidic flow velocities at the focal point, and transmission spectra discussed above, we
determined that an OPtIC microlens consisting of $9 \times 9$ array of nanoholes ($d = 150 \text{ nm}$ and a periodicity of $a_0 = 380 \text{ nm}$) with a center aperture of $d_c = 500 \text{ nm}$ is ideal for sorting of small size particles, such as exosomes. This microlens design is used in the rest of this paper.

High-intensity light sources, such as light emitting diodes (LEDs), are preferable to lasers for label-free size-based sorting applications. In this respect, NHA microlenses providing a focusing capability with minimal chromatic aberration are advantageous. We analyzed the effect of center nanofluidic channel on the broadband focusing characteristics of our plasmofluidic microlens for a wavelength range spanning from $600 \text{ nm}$ to $780 \text{ nm}$. Figure 3a shows that $f_D$ decreases monotonically with increasing wavelength. However, a particularly small focal length variation $\Delta z \approx 200 \text{ nm}$ is observed for the wavelength range $620 \text{ nm} < \lambda < 680 \text{ nm}$ corresponding to the (1,0) resonant transmission (EOT) peak. Outside the EOT spectral window, a significantly longer focal length is found at the transmission minimum $\lambda = 600 \text{ nm}$ (Fig. 3a) corresponding to the Wood's anomaly. We calculated focal spot size ($w$) along the focal plane denoted by the horizontal dashed lines in Fig. 3a. Within the (1,0) resonant transmission peak, only a slight difference is observed for the focal spot size. As shown in Fig. 3b, $w$ is $1.08 \mu m$, $1.12 \mu m$, $1.24 \mu m$ and $1.28 \mu m$ at $\lambda = 620 \text{ nm}$, $633 \text{ nm}$, $655 \text{ nm}$ and $680 \text{ nm}$, respectively. Similarly, light intensity profile along the OA presents minimal variations within the $620 \text{ nm} < \lambda < 680 \text{ nm}$ wavelength range (Fig. 3c). Hence, our analysis confirms that the (1,0) resonance transmission (EOT) peak provides a focusing behavior that is unaltered over a sufficiently-broad range of wavelengths.

The focal length of a finite-size opening that has identical dimensions to our plasmofluidic microlens can be calculated using the Rayleigh-Sommerfeld (R-S) formula:

$$ \frac{dI}{dz} = -2I_0 \frac{\pi \rho^2}{\lambda^2} \sin \left( \frac{\pi \rho n}{\lambda z} \right) = 0 $$

where $z$ is the distance above the lens, $I$ and $I_0$ are intensity and its peak values, respectively, whereas $\rho$ is the radius of the opening and $n$ is the refractive index of the surrounding medium. Our analysis shows that focal length $f_D$ of our plasmofluidic microlens (Fig. 3d, black curve) is in good agreement with the values obtained from R-S formula (Fig. 3d, grey dots) for the wavelength range $620 \text{ nm} < \lambda < 680 \text{ nm}$. Depth of focus (DoF) of our plasmofluidic microlens (shaded area bounded by the dashed lines) is shown in Fig. 3d. Even though the periodicity of nanohole array is broken with the incorporation of a nanofluidic channel ($d_c = 500 \text{ nm}$) at the center, our analysis...
confirms that our plasmo-fluidic microlens provides a well-defined focusing behavior with minimal chromatic aberration for the wavelength range of 620 nm < λ < 680 nm.

**Optical scattering, thermo-plasmonic and fluidic drag forces.** Optical scattering forces acting on a bioparticle can be decomposed to two orthogonal components: a radial gradient force ($F_g$) directed towards the beam axis and optical scattering force ($F_s$) in the direction of beam propagation against the fluidic flow. These forces can be expressed as:

$$F_{s,g} = \frac{2n_m P}{c} Q_{s,g}$$

(2)

where $P$ is the power of the incident light, $n_m$ is the refractive index of the medium, $c$ is the speed of light, and $Q_{s,g}$ is a dimensionless parameter defined for the scattering ($s$) and gradient ($g$) forces. $Q_{s,g}$ represents the efficiency of optical pressure transfer as a result of the light reflection/refraction at material interfaces. For simple beam profiles and symmetric geometries (i.e., a mildly focused Gaussian beam acting on a spherical particle), it is possible to calculate $Q_{s,g}$ analytically. For more complex beam profiles and small size particles with a diameter $a < 1 \mu m$, instead of the ray-optics model, optical forces must be calculated using Maxwell stress tensor (MST)\textsuperscript{28,29}:

$$T_{ij} = \varepsilon E_i E_j^* + \mu H_i H_j^* - \frac{1}{2} \delta_{ij} (\varepsilon |E|^2 + \mu |H|^2)$$

(3)

where $E$ and $H$ are the electric and magnetic field vectors, $\varepsilon$ and $\mu$ are the electric permittivity and magnetic permeability of the medium, and $\delta_{ij}$ is the Kronecker delta. Using FDTD simulations, MST allows us to obtain scattering and gradient forces acting on a particle for an arbitrarily shaped electromagnetic (EM) field distribution. Assuming a bounding box, small enough to contain only the particle of interest, the net optical force on the particle can be calculated as\textsuperscript{29}.
where $S$ is the surface of the bounding box and $\hat{n}_j$ is a unit vector along one of the principal axes.

For light intensities used here, electromagnetic heating of the OPtIC microlens can lead to large enough local temperature gradients inducing a buoyancy-driven convective flow away from microlens surface\textsuperscript{30,31}. A comprehensive discussion of heat induced fluid dynamics can be found elsewhere\textsuperscript{32,33}. Here, contribution of thermo-plasmonic effects is incorporated using finite element method (FEM). We first solve the electromagnetic wave equation for the electric field $E$ around the OPtIC microlens with a 500 nm-diameter center aperture\textsuperscript{34},

$$\nabla \times (\nabla \times E) - k_0^2 \varepsilon(\mathbf{r}) E = 0$$

where $k_0 = 2\pi/\lambda$ is the free-space wave number, and $\varepsilon(\mathbf{r})$ the position-dependent complex dielectric permittivity at the incident light wavelength. The calculated electric field distribution is used to obtain the heat source density $q(\mathbf{r}) = 0.5 \text{Re}[\mathbf{J} \cdot \mathbf{E}^*]$, where $\mathbf{J}$ is the induced current density in the microlens\textsuperscript{34} and the total heat power $Q = \int \int q(\mathbf{r}) \, d\mathbf{v}$. We then used coupled steady-state heat transfer and incompressible Navier-Stokes relations to calculate the temperature and thermo-plasmonic velocity field distributions

$$\nabla \cdot [\rho \kappa T(\mathbf{r}) \mathbf{u}(\mathbf{r}) - \kappa \nabla T(\mathbf{r})] = Q(\mathbf{r})$$

$$\rho \beta [\mathbf{u}(\mathbf{r}) \cdot \nabla \mathbf{u}(\mathbf{r}) + \nabla p(\mathbf{r}) - \eta \nabla^2 \mathbf{u}(\mathbf{r}) = \mathbf{F}$$

where $\nabla \cdot \mathbf{u} = 0$, $T(\mathbf{r}), \mathbf{u}(\mathbf{r})$, and $p(\mathbf{r})$ are the spatial temperature, fluid velocity, and pressure distributions, respectively. The material parameters for thermal conductivity $\kappa$, density $\rho$, specific heat capacity $c_p$, and kinematic viscosity $\eta$ are taken from ref.\textsuperscript{33}. The steady-state temperature distribution on the OPtIC microlens surface under 20 mW illumination at 633 nm is shown in Fig. S2a. Perpendicularly incident light transmitting diffractively through the center aperture is weakly coupled to the plasmonic excitations and dissipated within the metal film. Hence, a relatively low temperature difference from the ambient is observed within the center aperture region. Significantly higher local temperatures above the ambient are established outside the center aperture region. Through the center aperture is weakly coupled to the plasmonic excitations and dissipated within the metal film. These particles (a th $200 \text{ nm}$) experience stronger optical scattering and thermo-plasmonic convection forces that can push them against the fluidic flow lines ($F_{d}$). The robust size selective separation capability of our OPtIC platform is realized by harnessing the varying strength of optical scattering ($F_s$), thermo-plasmonic convection ($F_{tp}$), fluidic drag ($F_d$) and gravitational ($W$) forces based on size, refractive index and mass density of the bioparticles. The net forces ($F_{\text{net}} = F_s + F_{tp} + F_d + W$) acting on submicron particles (a $= 100 \text{ nm} - 1 \mu m$) at different locations along the OA ($z = 0 - 6 \mu m$) are shown in Fig. 4a.b. Here, a refractive index of 1.55 and a mass density of 1.05 g/cm$^3$ are assumed for the interrogated bioparticles. In Fig. 4a.b, the blue (red) region corresponds to physical conditions (particle sizes and spatial locations along the OA) leading to a net force towards (away from) the center aperture. For particles with a small diameter ($a_{th} < 200 \text{ nm}$), when 20 mW incident light ($\lambda = 633 \text{ nm}$) and a fluidic flow velocity of 1.3 m/s is used, optical scattering and thermo-plasmonic convention forces are weaker than the fluidic drag and gravitational forces ($F_d + W > F_s + F_{tp}$) at all locations along the OA (Fig. 4a, the vertical line on left). Hence, small diameter particles ($a_{th} < 200 \text{ nm}$) can readily follow the fluidic flow lines along the focal point and the high-intensity regions close to the center aperture. These particles ($a_{th} < 200 \text{ nm}$) make their way through the center aperture to the other side of the suspended OPtIC microlens and get separated from the mixture in the top channel (negative depletion). Under the same operating conditions, larger diameter particles (a $> 200 \text{ nm}$) experience stronger optical scattering and thermo-plasmonic convection forces that can push them against the fluidic flow lines ($F_s + W > F_d + F_{tp}$). These larger bioparticles ($a_{th} > 200 \text{ nm}$) are retained in the top channel (Fig. 4a). Threshold bioparticle diameter $a_{th}$ used for the size based fractionation can be readily tuned to a desired value by changing the light intensity and the fluidic flow velocity. In Fig. 4b, we show that the increased fluidic flow (velocity $= 3.0 \mu m/s$) shifts the threshold diameter ($a_{th}$) for size based sorting to 350 nm. By fine-tuning the relative contributions of the optical scattering, thermo-plasmonic convection and fluidic drag forces, microvesicles (up to 500 nm in diameter) can be selectively separated from larger bioparticles.
Minor differences in internal structure and chemical composition of bioparticles lead to effective refractive index variations that can be exploited for selective separation. A remarkable example of this is a recent study demonstrating that cells with single gene modifications can be distinguished based on their differences in chemical makeup using optical chromatography. Based on the same physical principles, our plasmofluidic platform demonstrating that cells with single gene modifications can be distinguished based on their differences in chemical makeup using optical chromatography.

Figure 4. The net force ($F_{\text{net}} = F_s + F_{tp} + F_d + W$) acting on submicron particles ($a = 100 \text{ nm} - 1 \text{ m}$) along the optical axis ($z = 0 - 6 \text{ m}$) for flow conditions corresponding to (a) $1.3 \mu \text{m/s}$ and (b) $3 \mu \text{m/s}$ flow rates at the focal point. We assumed that 20 mW incident light ($\lambda = 633 \text{ nm}$) and a fluidic flow velocity of $1.3 \mu \text{m/s}$ is used. Optical scattering ($F_s$), net fluidic drag ($F_d + F_{tp}$), gravitational ($W$) forces acting on a 200 nm exosomes-like bioparticle are shown as a function of particle refractive index ($n_e$) for (c) $1.9 \mu \text{m/s}$ flow rate and 100 mW incident power and (d) $0.95 \mu \text{m/s}$ flow rate and 20 mW incident power. The dashed vertical lines indicate the threshold particle refractive index ($n_{th}$) for which the net force acting the particles vanishes. Here, the lines are the first-order polynomial curve fits.

Virions, on the other hand, are a tight assembly of nucleic acids, proteins, and lipids; therefore have higher refractive indices ($n_{\text{virion}} \approx 1.48$) compared to exosomes ($n_{\text{exosome}} \approx 1.37 - 1.39$) that are closer to that of water ($n_{\text{water}} \approx 1.33$). Virions, on the other hand, are a tight assembly of nucleic acids, proteins, and lipids; therefore have higher refractive indices ($n_{\text{virion}} \approx 1.48$) compared to exosomes ($n_{\text{exosome}} \approx 1.37 - 1.39$). Accordingly, our platform enables us to use this refractive index difference to separate exosomes from similar size virions (Fig. 4c). Similar to the size-based separation (Fig. 4a,b), the threshold refractive index can be tuned to a desired value by adjusting the fluidic flow rate and light power (Fig. 4d). As demonstrated by Fig. 4d, at a relatively small flow rate ($0.95 \mu \text{m/s}$) and incident power (20 mW), the net force...
aperture through Coulter principle. Two electrodes added on the opposite sides of the microlens can readily read particle sizes by removing the larger size and higher refractive index particles using optical radiation pressure. The optofluidic platform employs a negative depletion mechanism enabling selective enrichment of exosomes in a single planar chip.

**Radial focusing of the bioparticles.** It is predicted that the instrumental fluctuations associated to variations in fluidic flow velocities can cause spatial dispersion of particles, deteriorating the size-based retention capability of conventional optical chromatography approaches. Instead of mildly focused Gaussian beams, our OPtIC microlenses use strongly focused light that can create large optical gradient forces in radial direction. As shown in Fig. 5b,c for 200 nm and 600 nm diameter particles, in addition to the optical gradient forces \( F_g \) (blue curve), the spatial dispersion of particles in our platform is inherently minimized by the fluidic drag \( F_{d,r} \) (black curve) and thermo-plasmonic convection \( F_{tp,r} \) (red curve) forces working together to align particles along the optical axis (\( r = 0 \)). Here, an incident light (\( \lambda = 633 \text{ nm} \)) power of 20 mW and a fluidic velocity of 1.3 \( \mu \text{m/s} \) are assumed. For distances up to \( \pm 1 \mu \text{m} \) away from the OA, the optical gradient \( F_g \) and fluidic drag \( F_{d,r} \) forces are minuscule and fluidic forces \( F_{tp,r} \) are dominant. Instead of mildly focused Gaussian beams, our OPtIC microlenses merge focusing capabilities of NHA plasmonic microlenses with nanofluidics. Our OPtIC microlens effortlessly realizes precise alignment of nanoparticles along the optical axis using radially focusing fluidic drag and optical gradient forces. We demonstrated that our plasmofluidic platform facilitates selective separation of nanoparticles below a threshold diameter, which can be tuned to a desired value using fluidic flow rate and the incident light power. Furthermore, we showed that our platform offers a robust separation capability for same size nanoparticles using differences in their chemical composition (refractive index). A major limitation in flow cytometry measurements is the diminished optical signals obtained from small size and lower refractive index nano-bioparticles, such as exosomes. Even with specialized flow cytometers equipped with high sensitivity detectors, the smallest detectable vesicles are typically larger than \( 200 \text{ nm} \) in diameter. Hence, majority of exosomes are overlooked in flow cytometry measurements, which is the most commonly used optical method in clinical and research laboratories. In contrast, our optofluidic platform employs a negative depletion mechanism enabling selective enrichment of exosomes in a readily manner by removing the larger size and higher refractive index particles using optical radiation pressure. Furthermore, our platform could be used for quantification of exosomes passing through the center nanohole aperture through Coulter principle. Two electrodes added on the opposite sides of the OPtIC microlens can detect brief changes (pulses) in the current that flows through the aperture, as the enriched exosomes transverse the enlarged center aperture. Previous studies have shown that resistive pulse sensing (RPS) technologies based
on Coulter principle in aperture format are capable of detecting bioparticles smaller than 100 nm. However, clogging of openings due to the larger particles has been a practical limitation when heterogeneous samples are analyzed. Our OPTIC microlens keeping larger diameter particles away from the nanopore aperture using only dielectrophoretic forces, when combined with RPS, could overcome these limitations and be used for selective sorting and detection of exosomes from heterogeneous samples. To achieve extended periods of operation and high-volume processing, one can incorporate a lateral microcross flow periodically removing the particle microfumes that build up above the focal point of the OPTIC microlens.

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**Author Contributions**

X.Z. and A.A.Y. conceived and designed the research, X.Z., A.C. and Y.L. performed research and conducted the numerical simulations, X.Z. and A.A.Y. analyzed the data. All authors contributed to writing of the manuscript.

**Additional Information**

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