Multi-access drug delivery network and stability

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Abstract: A novel design of a multi-drug delivery network and diagnosis using a molecular network is proposed. By using a pair of tweezers to generate the intense optical vortices within the PANDA ring resonator, the required molecules (drug volumes) can be trapped and moved dynamically within the molecular bus networks, in which the required delivery targets can be achieved within the network. The advantage of the proposed system is that the diagnostic method can be used within a tiny system (thin film device or circuit), which is available as an embedded device for diagnostic use in patients. In practice, the large molecular networks such as ring, star, and bus networks can be integrated to form a large drug delivery system. The channel spacing of the trapped volumes (molecules) within the bus molecular networks can be provided by using the appropriate free spectrum range, which is analyzed and discussed in the terms of crosstalk effects. In this work, crosstalk effects of about 0.1% are noted, which can be neglected and does not affect the network stability.

Keywords: drug delivery network, molecular networks, molecular diagnosis, neural system and network

Introduction

Human organs contain blood and tissue fluid. The heart pumps blood through the arteries and capillaries and returns to the heart via veins. By providing oxygen and nutrients to every cell of the body, all the cells are refreshed when molecules such as oxygen and nutrients move into tissue fluid from the blood. The blood circulation carries away waste products;1 each red blood cell 7–8.5 µm in diameter passes through the narrow capillaries smaller than 3 µm in diameter. Most capillaries range in diameter from 7 to 9 µm, and branch without changing in diameter. The circulation of blood through the human body is divided into two interlocking systems, venous and arterial. Together, they keep a dynamic interchange of blood moving to and from the heart and lungs.2 Several studies have been done to understand red blood cell transportation in the capillaries network,3 overall elasticity of the capillary system, apparent membrane viscosity, thickness of the double layer of electrical charges, adhesion of red blood cells in vascular to fabricate the same blood flow system,4 and pulmonary network via micro-fluidics system.5

Optical trapping was invented by Ashkin et al.6 It has emerged as a powerful tool with a wide range applications in biology, physics, engineering, and medicine.7 Optical trapping and manipulation of viruses, living cells, bacteria, and organelles without damage by laser radiation have been demonstrated.8–10 In medicine and the application of nanotechnology, a single red blood cell (RBC) deformability test has been performed by
optical trapping plastic in microfluidics chip and lab-on-a-chip for RBC transportation in capillary network to circulate oxygen and carbon dioxide throughout the human body. Optical trapping for manipulation of molecules in liquid core capillaries and its application to drug delivery has been reported by Suwanpayak et al, who used a PANDA ring resonator to form, transmit, and receive the microscopic volume of drug. The references of the practical devices are also given.

In the system proposed in this paper, the blood circulation system and pulmonary network can trap and transport (filter) drug from heart to capillaries. The required trapping tool sizes can be generated and formed for the specific blood circulation with oxygen and finally the clean blood can be sent to the destination via the through port. However, in practice, several sensors are required for environmental and blood quality control, which need to be explored. RBC transport in the capillary network is an indispensible element for this comprehensive model as well as lab-on-a-chip for RBC transport in capillary networks to circulate oxygen and carbon dioxide throughout the human body. This study investigated the use of two different-wavelength tweezers, molecular buffers, and bus networks to form the transported drug volume, especially for delivering and transporting large volumes of drug, suitable for a multi-drug delivery network such as molecular diagnostic networks, blood circulation networks, Alzheimer’s and Parkinson’s diagnosis, and molecular electronics. In addition, two different-wavelength tweezers were fed into the network to investigate molecular network stability. In practice, the multi-drug delivery networks could be used for large-scale drug delivery.

**Principle and method**

In theory, the trapping forces are exerted by the intensity gradients of highly focused light beams to trap and transport the microscopic volumes of matter. The optical forces are customarily defined by the relationship between optical scattering force and gradient force ($F_{grad}$). Furthermore, in the Rayleigh regime, the trapping forces decompose naturally into two components, since the electromagnetic field is uniform across the dielectric. Thus, the particles can be treated as induced point dipoles. Increasing the numerical aperture (NA) increases the gradient strength due to a decrease in focal spot size which can be formed within the tiny system, for instance, a nanoscale device (nanoring resonator). In this proposed system, the trapping force is produced by a dark soliton, in which the valley of the dark soliton is generated and controlled within the PANDA ring resonator by the control port signals. Figure 1 shows the output field $E_{out}$ at the through port. In the add/drop device, the nonlinear refractive index is ignored because it does not affect the system. The electric fields $E_0$ and $E_{out}$ are the fields circulating within the nanoring at the right and left side of the add/drop optical filter. To form the broad spectrum output, two nonlinear ring resonators are introduced to form a proposed structure called

$$0 \leq t < \tau$$
a PANDA ring. A panda is a well known Chinese bear, which was used to describe the polarization maintaining fiber core structure. In this work, the proposed ring resonator, called a PANDA ring, is a modified add/drop filter. By using the broad-spectrum output, the multitweezers can be generated and molecules can be trapped and transported.

The power output ($P_t$) at the through port is written as

$$P_t = |E_{n1}|^2. \quad (1)$$

The power output ($P_d$) at the drop port is

$$P_d = |E_{n2}|^2. \quad (2)$$

**Multi-access drug delivery network**

A molecular buffer needs to be included in the system. The molecular buffer plays an important role for storing or delaying atoms/molecules over a period of time, which gives enough time for operation.27-28 A molecular buffer is a new device, which is operated in the same way as a gas buffer.29 The polarizability of the particle is calculated by Equation (5). In this case, we assume that the spherical particle is polystyrene ($n = 1.5894$) and the liquid medium is water ($n = 1.33$). The optical power which is required to trap particles of a certain size/polarizability is 9.1 W (Figure 2). In simulation, the bright soliton with center wavelength 400 nm, peak power 1 W, and pulse width of 35 fs is fed into the system via the input port, where the coupling coefficients are $\kappa_a = 0.5$, $\kappa_i = 0.35$, $\kappa_j = 0.1$, and $\kappa_i = 0.35$. The ring radii are $R_{add} = 20 \mu m$, $R_R = R_L = 5 \mu m$. The evidence for the practical device with a radius of 2–3 $\mu m$ has been reported by Zhu et al.30 In this case, the dynamic tweezers (gradient fields) are in the form of bright solitons, Gaussian pulses, and dark solitons, which can be used to trap the required microscopic volume. In this investigation four tweezers with different center wavelengths are generated, whose dynamic movements can be seen in Figure 4, where Figure 4A represents tweezers with different sizes and wavelengths and Figure 4B represents tunable tweezers by coupling constant variation. The required drug volumes can be obtained by the drop port outputs.

In practice, the fabrication parameters can be easily controlled by the ring resonator radii instead of coupling constants. The important aspect of this system is that the tunable tweezers can be obtained by tuning (controlling) the add (control) port input signal, in which the required number of microscopic volumes (atom/photon/molecule) can be obtained at the drop/through ports, otherwise, they propagate within a PANDA ring without collapsing/decaying into the waveguide. In practice, the trapped drug molecules can be transported into the wavelength router via the through port, while the retrieved drug volumes are received via the drop port (connecting target). The advantage of the proposed system is that the transmitter and receiver can fabricate on-chip and alternatively can be operated by a single device. The magnitude of optical trapping force is in the pico Newton.
(pN) range, depending on the relative refractive index of particle.\(^3\) The particle radius located in the cavity decreases with the decrease in refractive indices compared with the host medium.\(^3\),\(^3\) The waveguide of the drug delivery system can be an optical waveguide with a liquid core which can trap the drug molecules smoothly within the network. By using the drug bus network, the trapped drug molecules can be transported to the required drug targets and the specific drug molecules can be obtained by using the molecular transceiver. To form the trapping tools, the PANDA ring resonator with four ports was used, as shown in Figure 1. First, the dark soliton is fed into the system via the input port. Second, the output trapping tools are transmitted into the throughput port and bus networks. Third, the required drug molecules are filtered and obtained via the drop ports. Finally, molecules are transported within the bus (ring) networks and drug routers, in which the control port is available for additional applications. The molecular trapping probe can be adjusted to select the drug molecule size of 80 nm or \(0.268 \times 10^{-3} \mu m^3\) per potential well (Figure 3A and B), but the fluidsics microscopic volume can be transported faster, which depends on the viscosity of media and particle.\(^3\) This molecular trapping probe can be used for drug molecule transport at the through port and networks, the parameters for which are given in the figure captions (Figure 3). The advantage of the proposed system is that it provides multiple access to the drug volumes and targets. Moreover, the use of mesh networks (combined networks) can also be realized, which can offer a large diagnosis area.

**Network stability calculation**

Several reports have shown that fluidics particles (drug volumes) can perform the realistic applications.\(^3\),\(^3\) The system proposed here shows that a tiny device in the form of thin film can be fabricated and used\(^3\) to integrate drug delivery network into the application area, as shown in Figure 3. Moreover, the use of the proposed system for the blood circulation network of artificial bone is also suitable for in situ surgery and neural and brain diagnosis. By using the design networks, the required trapped volumes can be transported within the network via the molecular buffer (storage) to the required destinations, for instance, the trapped tangle protein can be filtered via the add/drop filter before reaching the desired destinations. The throughput port \((E_{t1})\) output of add/drop filter is connected to the axon (axon terminal), then to neural cell and dendrite. The effective area of the waveguide is 2.01 \(\mu m^2\) \((r = 800 \text{ nm})\) and the outside diameter of the microtubule is 25 nm.\(^3\) Axon diameter at birth is 1 \(\mu m\), increasing through...

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**Figure 2** Results of the trapping tools. (A) wavelengths center 400 nm, (B) wavelengths center 450 nm. (C) and (D) are different tweezer separations. \(R_L = 2 \mu m\), \(R_i = R = 1 \mu m\). The coupling coefficients are \(\kappa_0 = 0.5\), \(\kappa_i = 0.35\), \(\kappa_L = 0.1\) and \(\kappa_i = 0.35\). The input power is 1 W.
childhood (7 years) to 12 µm and to 24 µm at adulthood. In Alzheimer’s diagnosis the optical tool is connected between the axon and the nerve cells and can be used to trap the tangle protein into the removal storage by add/drop filter (control port). The bus network design can also be used to trap the molecular motor to activate the information of neuronal cell at the same time. For better access, the coupling material is required to use as waveguide–axon coupling.

In operation, as “networks” are made up of add/drop filters (Figure 4), their performance depends on the add/drop filters. Micro- and nanowaveguides are gaining prominence in this field. Filters offer good stability and isolation between channels at moderate cost. The add/drop filters’ capability affects the network size. The maximum nodes of a network depend on the maximum amount of channels of add/drop filter. The popular dense wavelength division multiplexing (DWDM) component with many channels has been achieved in both laboratory and theoretical works. This means that multi-variable routers or networks with many ports can be built in future.

In this work, we propose the use of an optical network principle to estimate network stability. In general, the problems of a large network having the stability of a small network are insertion loss (IL) and crosstalk effect (FC). The IL reduces the efficient transmission distance. In popular communication networks, in which the signals pass through the router, insertion loss is usually 5 dB. According to the performance of present point-to-point transmission systems, we can build a required network over at least 50 km. Along with the development of DWDM technology, the insertion loss can be reduced to less than 0.5 dB. Therefore, the DWDM technology is promising for building large optical networks.

In order to test the proposed network principle, we use the DWDM technology to build a multi-access drug network for the treatment of Alzheimer’s disease. The network consists of 1000 nodes, each node representing a neuron. The input power is 1 W. The efficiency of the network is calculated as the ratio of the output power to the input power. The efficiency of the network is 95%.

Figure 3 Results of the trapping tools. (A) wavelengths center 400 nm, (B) wavelengths center 450 nm, (C) tweezers separation, (D) normalized tweezers, (E) multi-tweezers, and (F) normalized tweezers. \( r_{\text{add}} = 10 \mu m, R_1 = R_2 = 3 \mu m \). The coupling coefficients are \( \kappa_0 = 0.95, \kappa_1 = 0.5, \kappa_2 = 0.2 \) and \( \kappa_3 = 0.5 \). The input power is 1 W.
loss can be reduced to less than 1 dB in future.\(^{42}\) Then the multivariable network will cover more than 100 km with high capacity. The crosstalk effect is mainly due to a signal of co-channel interference and adjacent-channel interference. The crosstalk can be considered in terms of channel separability. For a network, crosstalk brings bit errors, so it must be reduced as low as possible.

IL and FC can be estimated as:

\[
IL = 10 \times \log \left( \frac{P_{in}}{P_{out}} \right) \quad (3)
\]

\[
FC_j(\lambda_i) = 10 \times \log \left[ \frac{P_j(\lambda_i)}{\overline{P}(\lambda_i)} \right] \quad (4)
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Here \(P_{in}\) and \(P_{out}\) are input and output of soliton, \(\overline{P}(\lambda_i)\) is output of soliton with wavelength \(\lambda_i\), and \(P_{in}\) is a function of the input power. We first assume that all input photons from any user are the same when they enter the router. Since one soliton can pass through two add/drop filters when it passes a router, the crosstalk versus efficient signals is given as:

\[
\frac{P(\lambda_i)}{P_{in}} = \frac{P(\lambda_i)}{P_{out}} \times \frac{P_{out}}{P_{in}} = 10^{\frac{(IL)(\lambda_i-IL)}{10}} \quad (5)
\]

\[
P_{out}/P_{in} = 10^{-IL/10} \quad (6)
\]

\[
\left[ \frac{P(\lambda_i)}{P_{in}} \right]^{2} \times \left[ \frac{P_{in}}{P_{out}} \right]^{2} = 10^{\frac{2(FC_j(\lambda_i))}{10}} \quad (7)
\]
Now we consider the situation in which the input soliton is not the same. A bad situation is that input photons which produce efficient signals pass through a device that has \( X \) dB insertion loss before passing through the router but those solitons which produce crosstalk do not. The ratio in Equation (7) will become \( 10^{\frac{X+2\pi FC(j)\lambda}{10}} \). If there are many inputs that produce crosstalk, the ratio must be

\[
\sum_{j=1}^{N} \left[ \frac{X+2\pi FC(j)\lambda}{10} \right]
\]

(8)

Here \( j \neq i \), where \( N \) is the number of channels (receiver nodes).

In present study, the system dimensions have been reduced to be micro-/nanoscale. From Figure 4, the coupling ratio of each coupling point is 50:50, ie, 3 dB coupling power. The molecular bus network has \( N = 4 \), \( FC(\lambda) < -4.26 \) dB (when \( j = i \pm 1 \)), \( X = -0.97 \)dB. The normalized input is 0.8; \( \lambda_i \) and \( \lambda_j \) equal 400 and 450 nm, respectively. Their ratio is less than 0.113%. Therefore the errors resulting from crosstalk are less than 1 (10%) and can be ignored. Along with the development of DWDM technology, crosstalk will be smaller and the performance of the molecular router or network will improve. Thus, we can easily build a feasible multi-variable network for multi-drug delivery applications, ie, for large networks.

Conclusions
We have proposed an interesting system that can be used for multi-drug delivery networks. The trapped drug molecules can move into the liquid core waveguide and networks by using optical tweezers, in which drug can be trapped, stored, and delivered via the molecular network. Such a system can also be used for large-scale molecular drug delivery network and diagnosis. The mesoscopic particle can be trapped and transported within the waveguide and network such as nanocarrier (polymeric nanoparticles, dendrimers) and lipid-based drug carriers.\(^{43,44}\) By using practical device parameters, such a proposed system can be fabricated and integrated into a practical thin film device. The trapping and movement of molecules in the system can be used for certain diagnostic purposes and to deliver small molecules to their target organ in the human body. Network stability was also calculated and it was found that the crosstalk effects due to the two wavelength-trapping drug molecules can be ignored. The proposed system can be used for long-distance transport of drug molecules in multi-drug delivery networks, in which drug delivery or molecular communication can be performed via the wavelength router and bus network, which will be available for large network systems (neural systems) in the near future. The proposed technique can be used in the new era of electronics and communications, where the use of molecules, DNA, genes, and atoms can have various applications within a tiny system.

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Disclosure
No conflicts of interest were declared in relation to this paper.

References


