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| **A. Biosensors**  Sensor: a device that detects and responds to input (stimulus) from its physical environment.  Transducer: a more general device for converting energy from a given form into a different form    Two-component system  –Biorecognition (**bioreceptor**): facilitate specific binding to or biochemical reaction with a target  – **Transduc**tion: converts a biological binding event to a measurable transduction signal (fluorescent read out, amperometric, potentiometric)  Two-fundamental focus:  Specificity: the analyte being measured  Sensitivity: the quantity of the analyte  *Sensitivity vs LOD (Limit of Detection) vs Selectivity*  i. Sensitivity: Ability to discriminate between small differences in analyte concentration at a particular concentration; to respond reliably and measurably to changes in analyte concentration  Slope = sensitivity = dy/dx  ii. LOD: Minimum concentration that can be detected at a known confidence limit  **LOD = 3s/m** (s for standard deviation, m for slope)  iii. Selectivity: distinguish analyte from other species in the sample  **B. Optical Biosensor**  1. Sensing Interface  1) Components: Light source, Sensing interface (transducer)， Optical detector  2) Properties of light:  i. travels as waves  speed: c = 3×108m/s  **frequency = ν**  **angular frequency = 2piν wave length = λ = c/ν**  **wave vector = k = 2pi/λ**  ii. light travels in straight lines  iii. reflection, absorption, transmission    Red light: longer wavelength, higher penetration  iv. Engergy of photon **E = hν = hc/λ**  h = 6.63 x 10-34 Joule second  Total Engergy = Nhν Photon momentum = p = h/λ  3) Refractive Index  the speed of light = c/n (n: refractive index)  Wavelength = λ0/n, λ0 is the wavelength in vacuum  4) Total internal Reflection    **Snell’s Law: n1sin (θ1) = n2 sin (θ2)**  **Critical Angle θc =sin‐1(n2/n1)**  ***Snell’s law does not work when θ1>θc***  Substitute θ2 = 90° to calculate the angle not enter 2  e.g: Fluorescence Microscopy, touch screen  5) **Evanescent Field (Many sensor based on)**      Many waveguide/optical fiber-based biosensors rely on evanescent field  2. Spectrometry  **Beer-Lambert law: T=I/I0=10^-αL=10^-εcL**  α: absorption coefficient ; L: solution’s path length(cm); ε: molar extinction coefficient; c: concentration(mol/L); I0: original light intensity; I: light intensity after sample  **Absorbance = A = -log10 (I/I0) = αL = εcL**  Sometimes, absorbance is expressed in “OD”  OD=1: 10 times attenuation (10% light passing through); OD=2: 100 times attenuation (1% light passing through); OD=: 1000 times attenuation (0.1%  light passing through)  3. **Fluorescence** Spectroscopy  1) Fluorescence excitation: absorption of light  Ads: High contrast, fluorescence can be incited by specific biological or physical process.    **dmin = dAiry**    Axial Resolution-Depth of focus    dtot: depth of field; λ: wavelength of illuminating light; NA: objective numerical aperture; M: lateral magnification  4) Photoacoustic Imaging (PAI)  Combine optical (high resolution) and ultrasound  (great imaging depth) advantages. Noninvasive / No X-ray exposure.  5. Hyperspectral Imaging  Spectroscopy: Shine light to the matter, light can be reflected, absorbed, and scattered.  When an incidental beam passes through a medium  which differ in refractive index (RI). If the medium has higher refractive index, the fraction of reflected  radiation will be higher.      Spectrometer measures reflectance, absorbance, etc.  Method: point-scan, line-scan(hyperspectral imaging)    Applications: remote sensing. Imaging skin cancer and border decisions, virus screening  6. Imaging Analysis  Image processing covers four main areas: Image formation, Visualization, Analysis of image, Management of the acquired information    7. Microfluidic Optical Biochips  Biochip: a miniaturized laboratory capable of  performing thousands of simultaneous biochemical  reactions.  1)Technologies: Sensing chemistry, Microarray or Microfluidics, Reader and Signal processing  2)ads: sample saving, faster analysis, integration  **D. Smart Systems**  Nano robot: The technology of creating machines or robots are close to the microscopic scale of a nanometer (10^−9 meters). | Fluorescence–ground state to singlet state&back.  Phosphorescence -ground state to triplet state&back  Both are radiative transitions, emission of photons from an electronically excited state  *Energy of emitted radiation is less than that of absorbed radiation because a part of energy is lost due to vibrational or collisional processes. Hence the emitted radiation has longer wavelength (less energy). This peak distance is called* ***stokes shift.***  2) Fluorescence lifetime (FLT): the time a fluorophore spends in the excited state before emitting a photon and returning to the ground state.      3) **Quantum Yield: Φ= kr/(kr+knr) = krτs**  **Where τs: the lifetime of S1 state. τs = 1/(kr+knr)**    Brightness= ε x Φ, molar extinction coefficient (ε)  Factors affects: Internal inversion, quenching, intersystem crossing rate, temperature, solvent, pH, Energy Gap Law.  4) Applications: Environmental Monitoring, Medical Diagnostics, DNA Squenching, GeneticAnalysis  5) Fluorescence Emission Process: Excitation of a fluorophore through the absorption of light energy; A transient excited lifetime with some loss of energy; Return of the fluorophore to its ground state, accompanied by the emission of light.  6) Selecting Filtering      *Excitation Filter Range*: a little higher than the initial wavelength to the peak wavelength of excitation filter  *Dichroic Mirror Wavelength:* the peak wavelength of excitation to the peak of emission filter  *Emission Filter Range:* the peak of emission filter to the end of emission filter  *Dichromatic mirror allows light of emission wavelength to pass through, while light of the excitation wavelength is reflected. So the line of it is low when the line of emission is high.*  Ratio of covered area by detector= a2/4πr2  (the area of sphere’s surface)  4. Raman Spectroscopy  *Based on scattering* of light; the sample is irradiated with a coherent source, typically a laser beam.  **• Photons are not absorbed**  1)types of scattering      Scattering: intrinsic molecular effect which provides another way to study energy levels  Blue sky effects: Lower wavelength light has more scattering  Rayleigh scattering (elastic): Photon energy does not change; leaves the molecule in the same state  Raman scattering (inelastic): Stokes lines are those in  **E. Electrical Biosensor**  Challenges: high sensitivity, selectivity, low power  1. Electrochemical Sensor    1)Types:  Potentiometric, Amperometric, Conductometric  **2) Nernst Equation**  The potential of a potentiometric electrochemical cell      Q: reaction quotient; n**:** the number of electrons exchanged in the half-cell reaction.; E° for each cell: standard hydrogen electrode (S.H.E.), strongest oxidizing agents higher; strongest reducing agents lower .  2. Amperometric biosensors  **Amperes x time = Coulombs**  **1 Faraday = 1 mole of electrons = 96,500 coulombs**  **1 mole= 6x10^23**  These sensors with fixed voltage  1) Glucose biosensor        3. Randle Circuits  a Randles circuit is an equivalent electrical circuit that consists of an active electrolyte resistance RS in series with the parallel combination of the double-layer capacitance Cdl and an impedance (Zw) of  a faradaic reaction.    4. Bioelectricity  1)Passive: To measurethe electric signals produced by the activity of living tissues  Electroencephalography(EEG, brain); Electrocardio-graphy (ECG, heart); electromyography (EMG, muscle)  2)Active: To study the effect of electric fields due to an external device on tissue  Deep brain stimulation (DBS); Functional stimulation (FES); Cardiac defibrillation (CDF)  **e ≅1.602 × 10^-19 C**  **1 C= e⁄ 1.602 × 10^-19 ≅ 6.241× 10^18e**  **1 A = 1 C/s 1 V = 1J/ 1C**  3) Biopotentials  Features: Threshold, All-or-none response, Intensity  **Vm = Vin − Vout < 0**    4) Five phases of Action Potential  Below Threshold- Resting potential; Depolarization; Action potential; Repolarization; Hyperpolarization  5. Bioelectric Measurement  To measure EMG: First, place two electrodes on the skin surface, e.g., one of them away from the muscle to be measured; Second, measure the difference between the potentials at the electrodes: Vd = Vl − V2; Third, we have an electronic circuit (amplifier) inside the acquisition system that amplifies Vd | which the photon has lost energy to the molecule, ν0–νt; Anti‐Stokes lines are those in which the photon has gained energy from the molecule, ν0 +νt  Raman spectroscopy studies the frequency change of  Light due to the interaction with matter  4. Optical Detector  1) Spectrometer, Monochromator, CCD, CMOS2) Dispersion of Grating: difference in the angle of  diffraction per unit change in wavelength      The smaller d, or the higher #groove density, the higher dispersion, higher spectral resolution  **3) Resolving Power or Resolution of Grating**  R = mN = λ/△λ  m = i: ith order diffraction pattern;△λ: smallest difference can be detected  5. Label Free Biosensors  *Analyte has different RI than that of environment (such as water, buffer, air); RI change before and after a binding event at sensor surface.*    1) Ads: No labeling involved; Direct sample measurement with minimal treatment or modification; No sample excitation; Low sample volumes needed; Potential rapid detection  2) Applications: Fiber and waveguide, Optical microcavity (Fabry‐Perot cavity, ring resonator); Surface plasmon resonance (SPR) (metal film and nanoparticle); Interferometer; Photonic crystals  3) Fiber Optics Biosensor  Ads: Many choices commercially available; Inexpensive; Easy surface functionalization; Compatible with catheters or endoscopes for in vivo biosensing; Sensing and light delivery in one device; Compact size; Multi-functional; Remote accessible  *Bottom Line for sensing:* ***You have to have access to the evanescent field***    Point: detect only one  Multiplexed: detect different location  Distributed: detect the same thing, but provide loca  Resonance wavelength change due to changes in radius ΔR or refractive index Δn:  Δλr /λr = ΔR/R + Δn/n    6. Resonator Biosensor  Longer interaction length → better detection limit  1) Disads: light passes the waveguide only once; Device size is large; large surface is needed, large sample quantity  2)Effective Length    λ-input wavelength; Pon-refractive index  3)Plasmon: The collective motion of electrons in  metal relative to ion background is called“plasmon”  Surface Plasmon Resonance Sensing    To measure ECG:          **x dB = 20lgx**  **Bode: y aixs for dB; x aixs for frequency**  6. Bio Amplifiers  Requirements: Biopotential amplifiers should have high input impedance i.e., greater than 10 MΩ; Output impedance of the amplifier should be low to drive any external load with minimal distortion; Gain of the amplifier is greater than x1000 as biopotentials are typically less than a millivolt; Safety: the amplifier should protect the organism being studied | •Fixed wavelength light, in a fan‐shaped form, is directed at the sensor surface and binding  events are detected as changes in the particular angle where SPR creates extinction of light.          4) Sensitivity to RI Change  with known refractive index    **C. Biomedical Imaging Technologies**  1. Applications  Magnetic Resonance Imaging, Computerised Tomography (High Resolution), X-Ray (Mostly common), Mammography, Position Emission Tomography, Ultrasound Imaging, optical imageing (No radiation, high sensitivity).  2. Photon interaction with biological tissue: Scatterd and reflected, and abosorbed, and transimitted      3. Oxygen Saturation and Concentration  Ads of Infrared: Non-invasive, non-radiaoactive, real-time functional imaging, portable, low cost  Estimating concentration by absorption coefficients:    𝜆1, 𝜆2: wavelengths; 𝜀𝑜𝑥, 𝜀𝑑e: Molar extinction coefficients of oxy- and deoxyhemoglobin; 𝐶𝑜𝑥, 𝐶𝑑e: Molar concentrations    Oxygenated hemoglobin absorbs more infrared light and allows more red light (660 nm) to pass through; Deoxygenated hemoglobin allows more infrared light (940nm) to pass through and absorbs  more red light  4. Optical Biospy (not to take tissue put to see)  1)Numerical aperture    The numerical aperture quantifies the capability of a lens to gather light: NA = n sinθ  n is the refractive index of the medium between the  objective lens and the specimen  θ is the half angle of the maximum light cone which the lens can collect.  2) Bright Field and Fluorescence Microscopy    Fluorescence: An excitation filter selects part of the electromagnetic spectrum for exciting the fluorescent materials in the specimen; Another filter is then utilized to separate the emitted light from that used for the excitation  **G. Response Robertic First Responder**  Monitor power: Bluetooth’s average energy is low but the peak power can be very high. Units can be damaged be current drop.  PPG: SPO\_2, heart contraction; capture potential signal, mechanical movement can cause noise, but the frequency is much lower than respiration so it will not generate electrical signal. Critical freq = 60  IMU: respiratory rate  ECG: pulse propagation, sensitive to movement  The **minimum distance between two ECG** electrodes should be more than 50mm: electrodes get electrical signal potential, hence if we wanta high enough big difference, we need big distance to allow signal go through enough distance so that there is a potential decay for ECG to capture.  **Best location to place implant**: implantation is under skin, we want somewhere that will not make objective pain, also we want the place where signals are detectable (ECG is usually 45°).    Redundancy Reason: none of the sensors can guarantee. Sensitive to noise (ambient lighting noise/ body movement)  Duty cycle: D = PW/T × 100%  PW: Pulse width; T: Total duty cycle period  Sleep/on: waking up the device can cost extra energy; it takes time to warm up to get reliable result since turning on needs to overcome inertia  With increasing frequency the Battery Life Time drops much faster than of continuous mode: Turning on has a higher power surge. So when the frequency is very high, doing nothing but waking up consumes energy. Waking up dominates the consumption  Drug Delivery: 1 revolution = 2 linear actuation  Mirror the figure:    Analog soft starter    Digital soft starter    DAC-based soft starter |