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Photo Pharmacology: A Door to Future Therapeutics

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ABSTRACT

For a drug to be effective, it should act precisely at its site of action. This indeed is a tiring work for drugs if they work alone. It takes help from light. This is a fascinating concept is what photo pharmacology is all about. Field of phototherapy was born with research work of Niels Finsen in the treatment of lupus vulgaris. Photodynamic therapy emerged at the same time with the works on combined effects of light and dyes. These branches of Photo medicine are in clinical use with few limitations. With the strong foundation of existing pharmacology and a new light technique came the field of so-called photo pharmacology. It aims at introducing photo switches into the drugs whose action can be switched on and off reversibly on irradiation. This review aims to give an overview on photo pharmacology and also its applications, current trends and future scope.

Keywords: Photo pharmacology, Phototherapy, Phototherapy in diseases

INTRODUCTION

Pharmacology deals with using drugs to treat disease. But for a drug to be effective target site selectivity is important. Most of the time, drugs are expressed not only at its target site but also in other sites leading to many adverse effects. Apart from this poor selectivity issues, Resistance is also emerging as another problem. These issues of poor drug selectivity, increased adverse effects and emerging resistance have literally compelled the researchers to explore new options which can add a completely new dimension to existing pharmacotherapy [1].

LIGHT IS THE NEW IDEA

From the era of bible, Curative ability of sunlight had been used to treat skin diseases. People of Egypt ingested "*ammi majus* "leaves for treatment of skin depigmentation. Later in 1850, Aaron Lerner was attracted to isolating the active component in these leaves which proved to be a photosensitizer when activated by sunlight. This was later followed by studies on safety and efficacy of these compounds in the treatment of vitiligo. All these pre historic trial and error experiments with light in medicine proved that light can potentially activate a substance by bringing out its therapeutic potential which led to the origin of a new field called Photo pharmacology (**Figure 1**) [2].



Figure 1. Light is the new idea. Source: Internet

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JOURNEY OF LIGHT IN MEDICINE

To understand this journey of light from Phototherapy to present era of photo pharmacology the following concepts play a role:

- Optogenetics is from Greek word "OPTOS" meaning Visible. Optogenetics undoubtedly has contributed in understanding biological process and is evolving as a major research tool but need for manipulation at genetic level is a major limitation.
- Photodynamic therapy is concerned with formation of cytotoxic ROS with photo switches that mediates transfer of Oxygen from triplet to singlet state upon irradiation. The fact that it can be used on sites where cell death should be evoked for treatment limits its use.
- Phototherapy deals with photochemical processes occurring in endogenous biomolecules. e.g., blue light induced isomerization of bilirubin for treatment of neonatal jaundice.

Both phototherapy and photodynamic therapy have issues in terms of poor drug selectivity and challenges in delivery of light. It has been seen that photo pharmacology is really successful in shining more light to the region of interest as in target selectivity.

Because main principle in photo pharmacology is not just targeting light to the site or altering at genetic level, it aims to alter the structure of the drug by introducing photo switchable moiety thereby achieving drug control. Main advantage when compared to other branches of photo medicine lies in its precision over target selectivity [2,3].

PROBLEMS IN CURRENT PHARMACOTHERAPY

Poor drug selectivity depends on the affinity of the drug for its targets. Cytotoxic agents like taxanes cause many adverse effects because these drugs act by impairing mitosis and this not only affects cancer cells but also healthy cells which causes all systemic side effects. To overcome this, targeted therapy should be achieved. Poor selectivity will lower the threshold level of toxicity thereby narrowing therapeutic window which leads to reducing the allowable dose. This results in not using the entire potential of the drug and probably we can conclude that many potential drug candidates which can improve current pharmacotherapy or even treat the untreatable are just discarded to the bin due to its poor selectivity.

Another major problem faced is resistance. This is of major concern in the field of antibiotics. The concept behind this idea of resistance and poor selectivity is that drug not only is selective for bacteria responsible for an infection in patient but also acts on the bacteria that are outside.

Effectiveness of a drug depends on both Pharmacokinetic and pharmacodynamics parameters. Pharmacokinetics deals

SciTech Central Inc. J Pharm Drug Res (JPDR) with how a drug is travelling around the body which is governed by ADME and can be evaluated based on the polarity, lipophilic nature and size of pharmacologically active compound. Dynamically these properties can be controlled by incorporation of a photo switch and photo irradiation. e.g., Polarity of the drug determines its distribution. Pharmacodynamics of a drug will define its drug-target interaction. Common drug targets are Ion channels, GPCR, Enzymes.

Main aim while designing any drug will be to achieve maximum affinity towards target. When photo switchable moiety is introduced into the drug, its configuration changes i.e.; drug switches between states of high and low affinity. So, at sites where drug action is not needed and to reduce adverse effects, drug can be possibly switched OFF and when action is required - switched ON [4].

CONCEPT OF PHOTO PHARMACOLOGY

Photo pharmacology uses light to modulate activity of drug as required. By inserting photo responsive switches to the parent drug changes its configuration when it is exposed to light of specific wavelength. These photo switches can bind by covalent or non-covalent interaction. Activation of these photo switches can happen either externally or internally. Photo switch is basically activated at a specific site with high spatiotemporal precision by using the specific wavelength hence rest of the sites in the body do not get exposed to harmful effects associated with the drug. This photo activation can be usually reversed spontaneously or by application of light of different wavelength [4].

PHOTO SWITCHES [5-7]

Definition: Compounds which absorbs photon and undergoes reversible change in both structure and property. Structure wise, they can belong to any type but should be compatible with presence of light and oxygen at cellular level.

Finding the Right Photo switch

Azobenzene molecule is usually favored as a photo switch over others.

Advantages: Once in a state of excitation, they undergo rapid photo isomerization on picosecond level. This prevents formation of toxic singlet oxygen which can kill the nearby cells. Azobenzenes are small molecules so they can be easily incorporated into LMW drug like molecule.

Disadvantages: Long term toxicity is a problem. When photo pharmacology is just applied like a research tool, this will not be a problem but when it reaches to the clinical trial level, this poses a serious concern.

WHY AZOBENZENES ARE TOXIC? (Figure 2)





Attaching Photo switch to Pharmacophore

This strategy is based on the fact that drug has high affinity for its target in one of the photoisomerization states.

- 1. PS can be incorporated directly into pharmacophore
- 2. Appended to pharmacophore

STEPS

- 1. Selection of suitable photo switch
- 2. Photo isomerization
- ✓ Consider suitable wavelength
- ✓ Consider reverse isomerization along with Photo isomerization (Usually photo switch that is OFF after stopping irradiation- Preferred)

Bistable Photo switch

- ON at a light of one wavelength & OFF at a light of different wavelength.
- Ideal photo switch will shuttle between the 2 states of configuration namely cis (active) and trans (inactive) states.

This is possible if at a particular wavelength, one isomer absorbs with a high molecular absorptivity where the other isomer is completely transparent.

3. Incorporate photo switch to the molecule of interest that brings about significant change in efficacy [4].

CRITERIA [8]

a) For targets (Table 1):

- Targeted by a drug that binds with high affinity and can modify its properties.
- Must be responsive to light induced changes in the structure of photo pharmacological agent.

• Should be accessible by light.

Table 1. Classification of targets in photo pharmacology [8].

Class	Accessibility	Example
1	Easily accessible	Skin
2	Accessible by endoscopy	GIT, Sinuses
3	Through skin without incision	Thyroid, Lymph nodes
4	Minor incision	Kidney, Intestine
5	Major incision intra operatively	Brain, Bone marrow

b) For photo pharmacological agents:

- Upon irradiation, increase in concentration of the active isomer should provide significant increase in response.
- Penetrating light must be of low toxicity
- Water soluble in both states
- Metabolically stable
- General toxicity with drug or its metabolites must be ruled out.

APPLICATIONS

A. Drugs and diseases

Alzheimer's disease [9]

One of the most difficult sites to irradiate with external light source will be the brain because we need invasive procedures to access it. But brain is usually the hub for many new drug developments since many disorders with complex pathologies and that are difficult to treat originate here. Photo pharmacology can help in studying different

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molecular targets in CNS but here we will try to focus on Alzheimer's disease.

So far Clinical trials of 105 drugs are in progress in all three phases for Alzheimer's disease where main target is to improve cognitive impairment and various other clinical parameters. But the overall rate of success in establishing a new drug is very low which can be attributed to complexity of disease process and understanding of enzymes and receptors involved. Poor target selectivity of these compounds also accounts for the low success rate.

Photo switchable inhibitor of Alzheimer's disease:

Hydrolysis of Ach is catalyzed by AchE. Reduction in Ach levels are found to be the reason for cognitive impairment in Alzheimer's disease so AchE inhibitors are the current treatment options now (**Table 2**).

Table 2. Photo switchable Inhibitor of Alzheimer's disease

 [9].

Molecule	Tacrine
Photo switch	Azobenzene
Active form	Trans (blue light)
Inactive form	Cis (UV light)

Antimicrobial Photo pharmacology [10]

Due to emerging resistance, need for localized delivery is increasing. Photo pharmacology serves this purpose by providing precise spatiotemporal control. In the field of antimicrobial photo pharmacology, mostly UV light is used for triggering the release of antibiotic which comes with its own risk of photo toxicity. Recently, Boron-dipyromethene sensitive at 470 nm was used as a photo switch and levofloxacin as a model drug. Here, visible light was used to cleave this drug complex (**Table 3**).

Table 3. Antimicrobial photo pharmacology [10].

Molecule	Levofloxacin	
Photo switch	BDP	
Light source	Visible light	
Advantage	Stable, nontoxic	
Photo analgesics [11] (Ta	ible 4)	
✓ Photo switchable opioids		
• Optical control over μ opioid receptor		
✓ Ta		
Table 4. Photo analgesics [11].		
4 .2 1 1		

Active moleculeFentanylPhoto switchAzobenzeneActive formTrans

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- Diabetes
- ✓ JB253
- ✓ Incretin switch

JB253 [12]

- Sulfonylurea is one of the important classes of drugs used in treatment of type 2 DM.
- M.O.A-binds to potassium channels (ATP sensitive)
- o Insulin secretagogues

Why Sulfonylurea is a target for photo pharmacology?

- Inexpensive option
- Safety profile well studied
- Most potent insulin secretagogue
- Designing photo switchable sulfonylurea (JB253) (Table 5)

Table 5. Sulfonylurea [12].

Active molecule	Sulfonylurea (glimepiride)
Photo switch	Azobenzene
Effect achieved	Optical control over insulin release

Incretin switch [13]

- Incretin mimetics
- M.O.A- Increases insulin secretion
- Promotes weight loss
- Problem- GI side effects
- Cyclic AMP-Increases insulin secretion
- Calcium and Increased cyclic AMP- Anti apoptotic effect (**Table 6**)

Table 6. Designing photo switchable incretin mimetic(LirAZO)[13].

Active molecule	GLP-1 receptor agonist	
Active molecule	(liraglutide)	
Photo switch	Azobenzene	
Trans form	Ca influx	
Cis form	Cyclic AMP generation	
D D		

- B. Receptors
- Nicotinic receptors [14]
- Location- Nervous system
- Application-Epilepsy, Parkinson's disorder, Alzheimer's disease

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- Target-Nicotinic Ach receptor
- Photo switch Azobenzene

For optical control

- Modify structure of the organic molecule by inserting photo switch
- Inactive caged molecule
- Cage is removed and active molecule released
- Acts on target
- Caged molecules targeting Nicotinic Ach receptor: NPE-carbachol, CNB-carbachol, RuBi-nicotine

Metabotropic glutamate receptors [15]

- GPCR
- Glutamate (excitatory neurotransmitter)
- Metabotropic glutamate receptors-3 groups (Table 7)
- Application-Treating anxiety, depression, Parkinson's disorder
- Photo switch used- azobenzene

Table 7. Metabotropic glutamate receptors [15].

Group I	Gq	Activates phospholipase- C	Calcium
Group II, III	Gi/Go	Inhibits adenylyl cyclase	Regulates ion channels

C. Ion Channels (Table 8) [16]

e.g., TRPC (Mammalian transient receptor potential channels)

Location- CVS, Brain

Function- Calcium entry

- CVS-target for Angiotensin II and Noradrenaline induced nuclear factor of activated T cells (NFAT)
- Involved in Cardiac stress responses like arrythmia, cardiac remodeling
- Overexpression-Promotes NFAT/Calcineurin pathway Promotes Hypertrophy and heart failure
- CNS- Calcium dependent degradation of purkinje cells-Impaired motor Coordination.

Table 8. Approach-Photochromic lipid approach [16].

Photo switch	Azobenzene
Lipid	DAG

D. Light to control protein [17]

Approach: Photo caging

Photo cage: Chemical group which is photo reactive that uses energy of photon and breaks a chemical bond placed usually at functional group in a biologically active molecule or amino acid of a protein by which activity is lost.

Upon irradiation- Photo cage removed-Release of a biologically active molecule.

E. Light to control enzymes [17,18]

Examples

1. HDAC2

Family- histone deacetylase

Function-Epigenetic regulation of gene expression

Overexpression - Cancer

Inhibition of HDAC2 kills tumor cells (HDAC2 inhibitor-VORINOSTAT)

2. NAMPT

- Function- NAD+ synthesis (salvage pathway)
- Increased in cancer cells
- NAMPT inhibitor-Reduction of NAD+ levels intracellularly-Apoptosis in cancer cell.
- APPROACH- Photo caging (Oxygen independent)
- Photo cage-Ruthenium
- Advantage-Visible light
- Since oxygen independent can be used to kill hypoxic tumor cells also. (Photo activated chemotherapy).
- F. Microtubules (Table 9) [19]
- Function-Mitosis
- Inhibit microtubule-Cancer cell death (Microtubule inhibitors)
- Problem- Side effects (peripheral neuropathy)
- Approach- Photo statins

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Table 9. Microtubules [19].

Active Molecule	Combrestatin A-4
Photo switch	Azobenzene
Active form	Cis (Voilet light)
Inactive form	Trans (Green light)
Method	C-C bond in Combrestatin A-4 with
in curiou	azobenzene

G. Tethered photo pharmacology [6]

- Variant of optogenetics
- Introduction of synthetic photo switch that changes effect of tethered ligand in a light dependent manner.
- Photo switching- Very fast so activation and deactivation within milliseconds

CHALLENGES

There is a compelling need for development of spontaneous photo switches since toxicity with existing azobenzenes also poses a major problem. Success of these light-based techniques will depend on a lot of factors like understanding more on existing photo switches involved and how they interact at the cellular and environmental level [1,4,20].

FUTURE OF PHOTO PHARMACOLOGY

Delivering photons to target poses a challenge in that they can get scattered at the tissue level and can be absorbed which may limit the penetration of light. With advances in optical techniques more precise control over spatiotemporal activity of drug can be achieved thereby improving safety and efficacy of these agents compared to their counterparts thereby paving way for clinical studies using photo pharmacological agents [6].

CONCLUSION

Photo pharmacology tries to create a difference in conventional pharmacotherapy to a greater extent by utilizing the mini- miracle photo switches more specifically azobenzenes and with the studies done so far, we can conclude that it is successful in dealing with problems like poor selectivity by employing techniques that have a high spatiotemporal precision. So far all is well when it is only happening in the research labs targeting receptors and enzymes but its real usefulness can be commented upon only when it targets the real population by clinical trials.

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