

Under the spotlight

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1 ***Under the spotlight: mechanisms of photobiomodulation concentrating***
2 ***on blue and green light.***

3 Short title: PBM with blue and green light.

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10
11 **Abstract**

12 Photobiomodulation (PBM) describes the application of light at wavelengths ranging from 400-
13 1100nm to promote tissue healing, reduce inflammation and promote analgesia. Traditionally, red
14 and near-infra red (NIR) light have been used therapeutically, however recent studies indicate that
15 other wavelengths within the visible spectrum could prove beneficial including blue and green light.
16 This review aims to evaluate the literature surrounding the potential therapeutic effects of PBM with
17 particular emphasis on the effects of blue and green light. In particular focus is on the possible
18 primary and secondary molecular mechanisms of PBM and also evaluation of the potential effective
19 parameters for application both *in vitro* and *in vivo*. Studies have reported that PBM affects an array
20 of molecular targets, including chromophores such as signalling molecules containing flavins and
21 porphyrins as well as components of the electron transport chain. However, secondary mechanisms
22 tend to converge on pathways induced by increases in reactive oxygen species (ROS) production.
23 Systematic evaluation of the literature indicated 72% of publications reported beneficial effects of
24 blue light and 75% reported therapeutic effects of green light. However, of the publications
25 evaluating the effects of green light, reporting of treatment parameters was uneven with 41% failing
26 to report irradiance (mW/cm²) and 44% failing to report radiant exposure (J/cm²). This review
27 highlights the potential of PBM to exert broad effects on a range of different chromophores within
28 the body, dependent upon the wavelength of light applied. Emphasis still remains on the need to
29 report exposure and treatment parameters, as this will enable direct comparison between different
30 studies and hence enable the determination of the full potential of PBM.

39 1 Introduction

40

41 The potential application of what is now known as Photobiomodulation (PBM) was first reported by
42 Endre Mester in 1967 at Semmelweis University, Budapest (1). Mester shaved the backs of mice
43 and shone a ruby red laser emitting a wavelength of 694nm on the backs of a group of mice in order
44 to investigate carcinogenicity. To his surprise, the hair on the backs of the irradiated mice grew back
45 faster compared with that of the non-irradiated control group. He called this phenomenon
46 'photobiostimulation' and to date (January 2019), over 6000 papers have been published regarding
47 the efficacy of PBM in treating a number of ailments by inducing analgesia (2), promoting wound
48 healing (3) and reducing inflammation (4).

49 PBM encompasses a broad range of different terminologies including low level laser/light
50 therapy (LLLT), cold laser therapy and phototherapy. Whilst the term PBM is the most recent
51 addition to this list, and is currently the preferred Medical Subject Heading Term (MeSH) which
52 encompasses both the stimulatory and inhibitory mechanisms involved, PBM is also often called
53 photobiomodulation therapy (PBMT) which further adds to the list of terms for the same therapy.
54 *Figure 1* gives an overview of PBM publications so far.

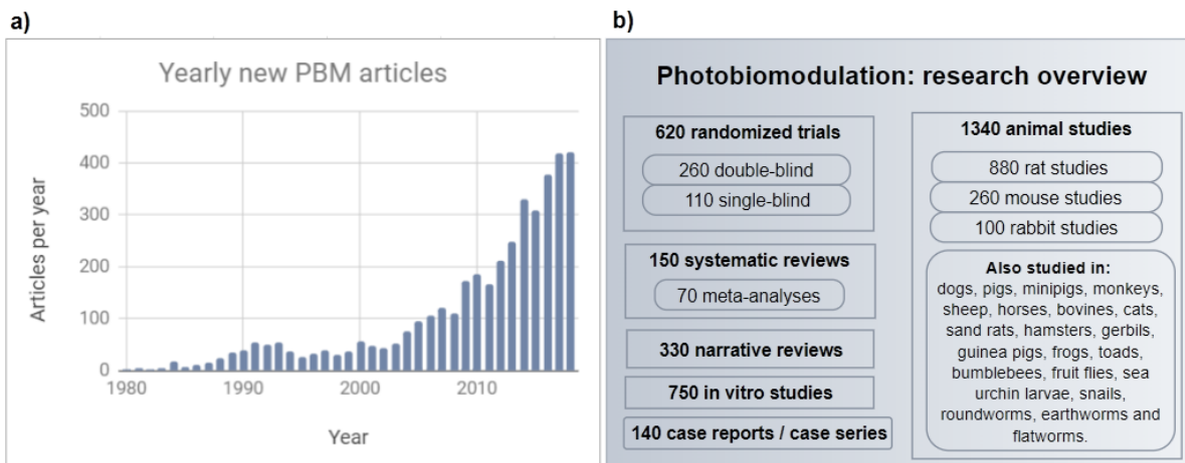


Figure 1: Preliminary PBM research overview based on a personal database of approximately 4,000 scientific articles related to PBM, compiled by manual PubMed and Google Scholar literature search using more than 80 different keywords (Supplementary file 1). (a) The amount of newly published PBM-related scientific articles has been increasing steadily during the 21st century, recently reaching a level of approximately 400 new articles per year. (b) The published research includes experimental in vitro studies and animal research. Also, a variety of randomized human trials and systematic reviews have been published so far.

55 Nonetheless, a growing number of observations suggests that specific wavelengths of
56 electromagnetic radiation spanning the visible to near infra-red spectrum (400-1100nm) could lead
57 to photo-physical and photochemical effects that can modulate major biological processes to
58 achieve therapeutic goals such as cellular proliferation, mitochondrial function and inflammatory
59 signalling (5) in various eukaryotic organisms, including humans. The majority of the literature
60 reports the beneficial therapeutic effects of red and near infrared light (red: ~600-750, NIR: ~750-
61 1100nm) in promoting tissue healing and reducing inflammation (6-14). Nevertheless, controversy
62 still surrounds the application of PBM in practice, due to the lack of knowledge concerning how PBM
63 elicits its molecular effects and also a poor understanding of photophysics and radiometric
64 parameters which affect repeatability and reliability (15). The importance of reporting treatment

65 parameters in a more consistent and reliable way has been emphasised in several articles and
66 guidance for reporting radiometric properties has previously been published (15, 16) to little or no
67 general avail (17-21). Indeed, those articles that have provided guidance for reporting radiometric
68 parameters have commonly recommended the consistent reporting of up to ten key radiometric
69 parameters (wavelength, power, irradiation time, beam area (at the skin or culture surface; this is
70 not necessarily the same size as the aperture), radiant energy, radiant exposure, pulse parameters,
71 number of treatments, interval between treatments and anatomical location) (22).

72 Whilst the majority of the literature supports the application of PBM using wavelengths
73 between 600-1100 nm, wavelengths <600 nm are less commonly researched or reported. The use of
74 blue light in particular (400-500 nm) is additionally surrounded by significant controversy relating to
75 the premise that the margin between 'safe' blue light and potentially damaging ultraviolet (UV) light
76 is not well defined.

77 UV light is divided into three discrete categories: UV-C (~100-280 nm), UV-B (~280-315 nm)
78 and UV-A (~315-400 nm) (23). A common misconception is that all UV radiation is associated with
79 DNA damage and mutagenesis (24). In fact, DNA damage is reportedly more efficient at UV-C and
80 UV-B wavelengths with a peak absorption at 254 nm which corresponds to absorption by one of the
81 nucleotide bases of DNA known as thymine, resulting in the formation of thymine dimers and
82 rendering the DNA molecule inactive and unable to replicate. UV-A radiation on the other hand has
83 a poor efficiency in inducing DNA damage, because it is not absorbed by native DNA or any of its
84 bases. However, like red and NIR wavelengths, UV-A wavelengths are able to generate singlet
85 oxygen (reactive oxygen species, ROS), and if the concentration of these radicals is in sufficient
86 quantity, they can damage DNA (25). However, ROS in small quantities can be beneficial to cells and
87 is commonly associated with proposed mechanisms of PBM (26). Indeed, the production of ROS is
88 likely to be influenced by radiometric parameters, namely, wavelength, irradiance, dose and the
89 number of photons delivered and again highlighting the importance of these parameters.
90 Nonetheless, the use of wavelengths in PBM at 400 nm or lower should be utilised in practice with
91 extreme caution.

92 In addition, another key caveat regarding the use of blue light for PBM is the low
93 penetration depth of blue light through tissue compared with that of red or NIR light (27). Whilst
94 blue light is cited to possess a penetration depth corresponding to an intensity decrease by 1/e (or
95 approximately 63%) at 1mm, NIR light has a penetration depth of up to 5mm through tissue (28).
96 However, there is a body of growing evidence supporting the use of PBM using blue light to reduce
97 inflammation in superficial tissues (29) and promote wound healing (30), as well as being able to
98 limit bacterial growth (31). Similarly, wavelengths within the green section of the visible spectrum
99 (495-570 nm) have also gathered considerable interest. Published reports have indicated PBM
100 effects for green light ranging from improved cellulite appearance (32) to reduced tissue swelling
101 (33).

102 This review aims to evaluate the main primary and secondary mechanisms involved in light
103 transduction in particular with blue and green wavelength of light. Secondly we focus on evaluation
104 of current literature regarding the therapeutic efficacy of blue and green PBM.

105

106

107

108

109 2 Primary mechanisms of PBM

110

111 According to the Grotthuss–Draper law, commonly termed “the First Law of Photochemistry”,
112 photochemical reactions are dependent on the absorption of light by a system. Subsequently in this
113 section, we provide a review of the literature of the most often proposed cellular photoacceptors
114 (chromophores) that are reported to mediate the biological effects in PBM. We cover in particular
115 the possible photoacceptors responsible for the transduction of blue and green wavelengths of light.

116

117 2.1 Cytochrome c oxidase

118

119 It has been proposed that PBM acts directly on the electron transport chain located in the
120 mitochondrial membrane, specifically on the enzyme cytochrome c oxidase (CCO), also known as
121 complex IV (34). The electron transport chain is comprised of five complexes: complex I (NADH-CoQ
122 reductase), complex II (succinate dehydrogenase), complex III (cytochrome c reductase), cytochrome
123 c oxidase and complex V (ATP synthase). Electrons are passed systematically down the chain of these
124 complexes in order to generate a proton gradient to provide the activation energy for ATP synthase
125 to catalyse the production of ATP (35). CCO is responsible for the conversion of molecular oxygen
126 (O_2) to two molecules of water (H_2O). CCO contains two copper centres (Cu_A , Cu_B) and two hemes
127 (cytochrome a, cytochrome a_3), which are involved in redox reactions within the enzyme.

128 The most widely accepted explanation for the beneficial photobiological effects of red and
129 near-infrared light has been the “CCO theory” largely established by Tiina Karu in the 1990s. It posits
130 that the light-cell interaction responsible for the observed PBM effects occurs initially at the redox-
131 active copper atoms of CCO complex in the mitochondrial electron transport chain (36-39). The CCO
132 theory was based on Karu’s earlier findings in the 1980s, which showed that the position of peaks in
133 the action spectrum measured for a variety of light-induced cellular changes (including DNA
134 synthesis, RNA synthesis and cell attachment) were practically identical. These findings suggested
135 that a universal cellular photoacceptor could be capable of absorbing those specific wavelengths and
136 producing cellular changes affecting multiple cellular compartments. The observed peaks in the
137 action spectrum were located within the blue (404 nm), red (620 and 680 nm) and near-infrared
138 (760 and 820 nm) parts of the electromagnetic spectrum (36).

139 Various *in vitro* and *in vivo* studies have observed effects related to increased mitochondrial
140 activity, including increased ATP levels, ROS levels, and mitochondrial membrane potential following
141 irradiation. Interestingly, the time it takes for these effects to become evident varies from minutes
142 to hours depending upon the experimental settings (40-42). Effects on mitochondrial function have
143 also been demonstrated in animal (43, 44).

144 Interestingly there remains no clear understanding, however, of the exact events that occur
145 within the electron transport chain or the enzyme CCO during light absorption to produce these
146 effects. A multitude of hypotheses have been proposed, including photodissociation of nitric oxide
147 (NO), changes in CCO redox properties with acceleration of electron transfer, superoxide generation
148 and biochemical changes related to transient heating of irradiated photoacceptors (45). It has also
149 been suggested that cytochrome c oxidation by CCO might be catalysed by red light irradiation (46).
150 However, a later replication study failed to confirm this effect, and also raised doubts indicating that
151 the initial positive results could have been experimental artefacts due to lack of detergent used for
152 the CCO solubilization (47). An alternative explanation for the observed mitochondrial effects could

153 also be an increased efficiency of CCO proton pumping (48). Regardless, the very limited amount of
154 observations allows no firm conclusions to be made on the subject.

155 The hypothesis of NO photodissociation from CCO is relatively popular, and based on the
156 understanding of the reversible inhibitory effects of NO on CCO (49, 50). There is some evidence
157 suggesting that light can attenuate the mitochondrial inhibitory effects of NO (51) and some
158 suggesting that NO can also inhibit the cellular effects of light (52). Light has been shown to increase
159 NO levels in cells and blood (53). However, the evidence is not completely consistent. One
160 experiment failed to demonstrate the protective effect of red light against NO-induced inhibition of
161 mitochondrial respiration, but demonstrated partial protective effects with blue light (442 nm) (54).
162 Another experiment with blue light (430 nm) recovered the mitochondrial function that had been
163 inhibited by nitric oxide at the levels generated under septic conditions (55). Hence, demonstrating
164 wavelengths outside the red and NIR range could be effective in modulating mitochondrial activity.

165

166 2.2 Opsins

167 Opsins are G-protein coupled receptors that have gained considerable interest in phototherapy
168 research due to their excitation by blue or green light (56) (see *Figure 2*). Opsins can be divided into
169 subcategories dependent upon the location they are expressed.

170 Opsin 1 (OPN1) and 2 (OPN2) expression is localised to the retina in the eye. OPN1 is expressed
171 by cone cells, photoreceptors within the eye that recognise coloured light and can be subdivided
172 into three types: OPN1 short wavelength (OPN1-SW), OPN1 medium wavelength (OPN1-MW) and
173 OPN1 long wavelength (OPN1-LW). Conversely, OPN2 (rhodopsin) is expressed by rod
174 photoreceptors, cells that recognise dim light and are important in peripheral vision (17). Three
175 further opsins are expressed within the human body including OPN3 (encephalopsin), OPN4
176 (melanopsin) and OPN5 (neuropsin) all of which exhibit an absorption spectrum ranging between
177 380-496 nm. Notably, OPN expression has been detected throughout the body with the expression
178 of OPN2, OPN3 and OPN5 being found in epidermal skin (57) and the expression of OPN3 in the
179 brain (58).

180 A number of publications have explored the role of OPNs in blue light mediated PBM
181 signalling both *in vitro* and *in vivo*. For example, *Regazetti et al.* (44) explored the effects of
182 irradiation at 415 nm (50 J/cm²) or 465 nm (62.5 J/cm²) on the regulation of pigmentation through
183 OPN signalling. The authors concluded that OPN3 could provide a novel target for regulating
184 melanogenesis (59). *Ortiz et al.* [45] also explored the effects of irradiation at 400 nm or 460 nm on
185 the influence of signalling through OPN3 and OPN4 on pulmonary vaso-relaxation (60). The authors
186 concluded that blue light induced vaso-relaxation and reduced arterial relaxation, through OPN3 and
187 OPN4 signalling. A series of further publications have also evaluated the role of blue light in
188 influencing opsin signalling to regenerate visual pigments (61) and promote hair regrowth (56).
189 Notably, whilst there is a wealth of literature supporting the idea that opsin signalling influences
190 responses both *in vitro* and *in vivo*, the molecular and cellular mechanisms of this signalling pathway
191 are yet to be fully elucidated.

192 Current literature indicates that different opsins are coupled to different subtypes of G-
193 proteins and hence induce different signalling pathways. For example OPN4 is coupled to Gq
194 (activates the phospholipase C pathway) whilst other opsins (OPN1, OPN2, OPN3 and OPN5) are
195 coupled to Go (inhibits adenylate cyclase), Gi (inhibits adenylate cyclase), Gt (transducing, activates
196 phosphodiesterase 6) and Gs (activates adenylate cyclase) (62-64).

197 It is proposed that one downstream target of opsins is transient receptor potential (TRP)
198 channels, particularly the TRPV1 subtype (capsaicin receptor), which has been cited to be activated
199 by light (65). TRP channels are ligand-gated ion channels. When a stimulus is applied, the TRP
200 channel opens and this enables a flood of calcium (Ca^{2+}) ions into the cytoplasm of the cell. In turn,
201 Ca^{2+} then induces the activity of calcium/calmodulin dependent kinase II (CAMKII), which in turn
202 induces the phosphorylation of the transcription factor, cAMP response element-binding protein
203 (CREB) located in the nucleus. In turn, CREB induces a series of changes in gene transcription
204 ultimately proposed to lead to some of the beneficial effects of PBM, seen both *in vitro* and *in vivo*
205 (59, 64). *Figure 2* highlights the current proposed molecular mechanisms relating to how blue and
206 green light PBM triggers opsin signalling. It has also been shown that increased activity of TRP
207 channels induces ROS generation (66) and thus the activation of the Ras pathway, a key pathway
208 involved in the modulation of the activity of small GTPases, ultimately leading to the modulation of
209 calcium signalling and apoptosis (67). Hence, these mechanisms may explain current findings which
210 indicate that blue light induces significant increases in ROS production (19, 68-72). However,
211 evidence also suggests this may be due to the effects of blue light on mitochondrial activity, inducing
212 increases in ROS production as a result of the stimulation of the electron transport chain (73).
213 Therefore, whilst there is a wealth of literature suggesting that blue light induces the activity of
214 opsins and TRP channels, the pathway that links these two complexes is yet to be elucidated.
215 Therefore, further work is required to determine the molecular mechanisms involved.

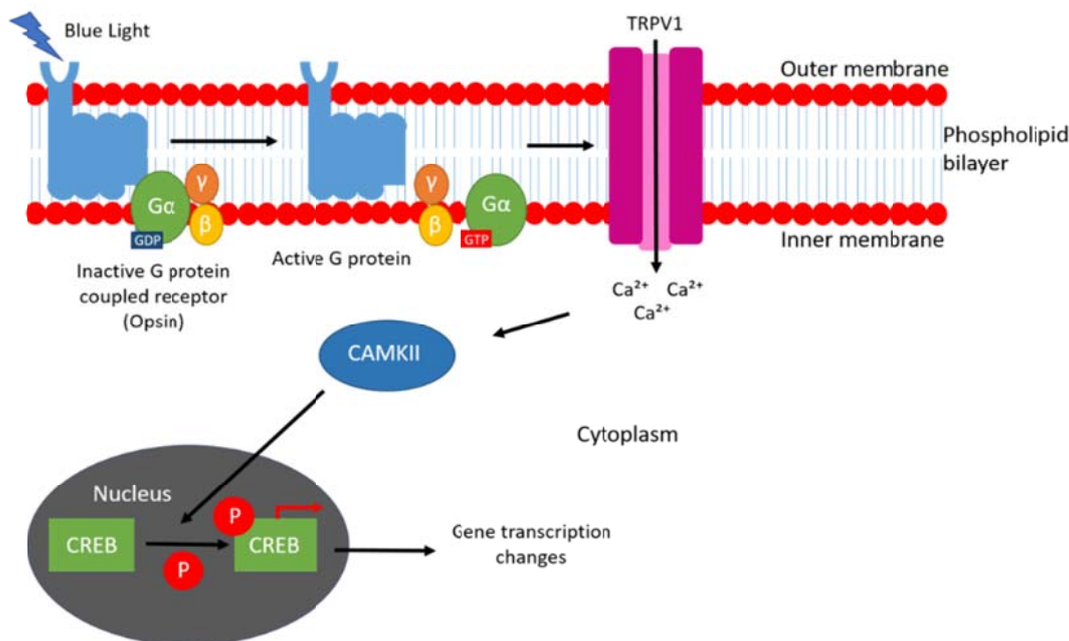


Figure 2: Possible molecular mechanism of blue light PBM in which an opsin receptor is activated by blue light, which induces a conformational change in the cis-retinaldehyde cofactor, allowing it to act as a guanine nucleotide exchange factor. This then enables the dissociation of guanosine diphosphate (GDP) from subunit $G\alpha$ of the associated G protein and the binding of guanosine triphosphate (GTP). In turn this provides the activation energy to enable $G\alpha$ to dissociate from $G\beta$ and $G\gamma$ (the other subunits of the G protein) and enables signalling of $G\alpha$ through a series of pathways including the cAMP and phosphatidylinositol pathways. In turn, signalling through these pathways is understood to induce the downstream activity of transient receptor protein (TRP) channels including the capsaicin receptor (TRPV1), which causes a flood of calcium ions into the intracellular space, resulting in the activation of calcium/calmodulin dependent protein kinase-II (CAMKII) and thus the phosphorylation of CREB (a transcription factor). CREB then induces a series of transcriptional events.

217 2.3 Flavins and flavoproteins

218

219 Blue light (400-500 nm) is known to excite flavins and flavoproteins including flavin mononucleotide
 220 (FMN) and flavin adenine dinucleotide (FAD) (74). A well-characterised family of flavin containing
 221 complexes is called "cryptochromes" (75). Notably, cryptochromes have been widely documented to
 222 absorb blue light (76) and are proposed to be involved in the regulation of the circadian rhythm in
 223 mammals (77). Notably, FMN is also found within complex I of the electron transport chain and it is
 224 proposed that blue light provides the activation energy for FMN to catalyse the reduction of oxygen
 225 (O_2) to superoxide ($O_2^{\cdot-}$)(78). Hence, blue light is understood to induce increases in the levels of
 226 circulating ROS (79). Complex II is also a flavin (contains $FADH_2$) containing cytochrome (80), and also
 227 absorbs blue light (81). Hence, it is plausible that like red and NIR light, blue light could affect
 228 mitochondrial activity. Indeed, *Serrage et al* demonstrated that blue light (400-450 nm, $5.76 J/cm^2$)
 229 was as effective in inducing increases in mitochondrial activity as NIR light (810 nm, $5.76 J/cm^2$) (82).
 230 However, further work is required to validate this hypothesis to determine whether blue light can
 231 modulate the activity of flavin containing complexes of the ETC.

232

233 2.4 Porphyrins

234

235 Porphyrins, a group of heterocyclic organic compounds found complexed to proteins ranging from
236 haemoglobin (83), to cytochrome p450 enzymes (84), to complex IV of the electron transport chain
237 (CCO) (85) are known to possess a typical Soret band at 400-420nm and hence possess the ability to
238 absorb blue light (86). Blue light of wavelengths between 400-415nm induces the π to π^* transition
239 in porphyrin rings (87). Wavelengths between 400-420nm could oxidise porphyrin containing heme
240 groups (found within complex IV), whilst a wavelength of 450 nm could induce CuB (a component of
241 complex IV) reduction hence inducing complex IV oxidation or reduction respectively (88).

242 When evaluating the influence of PBM on mitochondrial electron transport chain activity,
243 Evgeny et al concluded that blue light application (442 nm, 30 mW/cm², 3 J/cm²) induced significant
244 increases in complex IV activity and cell metabolic activity, compared to NO which inhibited cell
245 responses (54). Also, Ankiri et al reported that complex IV possesses a maximal absorption at 410 nm
246 and hence this could be due to porphyrins contained within the complex (89). Similarly, Del Olmo-
247 Aguado et al evaluated the effects of blue light on retinal ganglion cell mitochondrial activity. The
248 authors concluded that blue light upregulated the activities of complexes III, IV and V of the electron
249 transport chain, but also induced significant reduction in cell viability, and induced apoptosis (90).
250 These data indicated a possible role of blue light in affecting porphyrin-containing complexes of the
251 electron transport chain.

252 Cytochrome p450s (CYPs) are also porphyrin-containing complexes that have gained interest
253 in phototherapy research, as their activation by blue light has been cited (91). CYPs are a family of
254 proteins that contain heme and are vital for drug metabolism. The p450 element of the cytochrome
255 refers to the protein absorption spectra, since CYPs exhibit a maximal absorption peak at 450 nm
256 when bound to carbon monoxide (92). CYPs are membrane bound proteins and can be located
257 either in the endoplasmic reticulum or the inner mitochondrial membrane. Mitochondrially-located
258 CYPs including cytochrome P450 reductase, transfer electrons from nicotinamide adenine
259 dinucleotide phosphate (NAPDH) and thus could play a role in ETC activity (93).

260 Interestingly, Becker et al evaluated the effects of blue LED irradiation (453 nm, 23 mW/cm²,
261 41.4 J/cm²) on the proliferation and gene expression of keratinocytes, and found that irradiation
262 induced a decrease in cell proliferation. However, the authors also reported that blue light induced
263 significant increases in the transcription of electron transport chain-related genes, cytochrome P450-
264 related genes and also genes relevant to steroid hormone synthesis (94). Becker and colleagues also
265 reported that genes relevant to inflammation were significantly down-regulated due to this
266 exposure, and proposed that this may be due to the induction of steroid hormone biosynthesis via
267 the CYPs pathway. Hence, these data provide an additional hypothesis as to how blue light PBM
268 could affect cellular signalling.

269

270 2.5 Nitric oxide (NO)-containing compounds and nitrite reductases

271

272 In addition to the NO photodissociation hypothesis (see Section 2.1), there is some literature
273 suggesting that light-mediated effects are related to the synthesis combined with, or without, the
274 release of NO due to light exposure.

275 Firstly, CCO has also been shown to function as a nitrite reductase, thus being able to produce NO
276 locally in the mitochondria. This nitrite-dependent NO synthesis in isolated mitochondria has been
277 demonstrated to increase by yellow light (590 nm), without any concomitant increase in
278 mitochondrial oxygen consumption (100). These data suggest that light-induced cellular effects do
279 not necessarily have to be coupled with changes in mitochondrial respiration.

280 Secondly, there is also evidence that certain wavelengths of light can induce the release of NO
281 from photolabile sources of stored NO, such as nitrosyl hemoglobin (HbNO), nitrosyl myoglobin
282 (MbNO), S-nitrosothiols (RSNO) or dinitrosyl iron complexes (DNIC). This effect is reportedly much
283 greater with red light (670 nm) compared with some longer wavelengths that have been examined,
284 including 740 nm and 830 nm (101, 102). NO release from some S-nitrosothiols (RSNOs) has also
285 been demonstrated with ultraviolet (340 nm) and green (545 nm) wavelengths of light (103). Blue
286 light (420-453 nm) has been shown to be capable of eliciting NO release from S-nitrosoalbumin
287 (SNO-Alb), HbNO and aqueous nitrite solutions (104) (105).

288

289 3 Secondary mechanisms of PBM

290

291 From evaluation of the possible hypotheses of the primary mechanisms of PBM, it is apparent that
292 several pathways converge on the induction of the same signalling molecules; i.e. ROS. Therefore,
293 this section of the review evaluates the possible effects of PBM on ROS-related pathways. It is
294 prudent to highlight this review evaluates the effects of PBM on a number of downstream targets.
295 PBM may modulate a variable number of these targets, dependent upon the dose of light used, the
296 wavelength employed and also the *in vitro/in vivo* model light is applied to. The same is true for the
297 primary mechanisms of PBM, illustrated in section 2.

298

299 3.1 Nuclear factor kappa-light-chain-enhancer of activated B cells (NFκB)

300

301 ROS production instigates a signalling cascade ultimately leading to the phosphorylation of IκB, an
302 inhibitor of the pro-inflammatory transcription factor NFκB. In its inactive state IκB is bound to NFκB
303 in the cytoplasm, however, once phosphorylated, IκB dissociates from NFκB and is targeted to the
304 proteasome for degradation. This then allows the translocation of free NFκB to the nucleus binding
305 to DNA, and initiation of a series of gene transcription changes, mRNA production and potential
306 downstream expression of key cytokines, chemokines and growth factors including interleukin-8 (IL-
307 8), IL-6 and vascular endothelial growth factor (VEGF (106-109)).

308 A number of authors report the modulation of NFκB by light. For example, Chen et al
309 reported that irradiation at 810 nm and radiant exposure of 0.003 J/cm² induced the activation of
310 NFκB through increased ROS production induced by light (40). Similarly, Curra et al evaluated the
311 effects of a 660 nm diode laser on NFκB protein levels using an *in vivo* hamster model of oral
312 mucositis (110). The authors concluded that PBM reduced disease severity through the activation of
313 the NFκB pathway. Conversely, PBM reportedly may reduce NFκB activation and subsequently
314 reduce the expression of pro-inflammatory mediators in several diseases (111). Interestingly, de
315 Farias Gabriel also reported that application of 660nm (4J/cm²) modulated NFκB activation leading
316 to keratinocyte migration, resulting in improved wound healing in a rat model for oral epithelial

317 wound healing (112). Hence, modulation of NFκB not only affects pathways related to inflammation
318 but also those influencing wound healing.

319 Another gene directly regulated by NFκB activation is cyclooxygenase-2 (COX-2). Its main
320 role is to catalyse the conversion of arachidonic acid to prostaglandins including PGE₂ (113). PGE₂
321 has then been reported to be involved in the activation of a variety of pathways including cyclic
322 adenosine monophosphate/protein kinase A (cAMP/PKA) signalling (114, 115).

323 To stimulate activity of the cAMP pathway, PGE₂ binds to prostaglandin E₂ receptor 4 (EP4).
324 EP4 is a GPCR coupled to a stimulatory G protein (Gs). On binding, PGE₂ induces a conformational
325 change activating Gs, which then activate adenylyl cyclase to catalyse the conversion of ATP (a
326 second molecule whose production is increased by PBM) to cAMP (116). cAMP then induces the
327 activation of protein kinase A (PKA) leading to the phosphorylation of transcription factors including
328 CREB (117). Several authors have reported the effects of PBM on NFκB induced signalling. Lim et al
329 reported that irradiation at 635 nm modulated both COX2 and PGE₂ protein expression (118).
330 Current literature also indicates the effects of PBM on a series of signalling proteins/molecules
331 implicated in this pathway, including CREB (119).

332 Another key molecule modulated by NFκB signalling is VEGF, a growth factor central to the
333 promotion of angiogenic events (120, 121). Literature reports indicate that activation of EP4 induces
334 the upregulation of the expression of VEGF and several authors have reported the effects of PBM on
335 VEGF expression and activity. Tim et al concluded that 830 nm irradiation of male Wistar rats with
336 induced bone defects induced significant increases in COX2 and VEGF expression (122), and das
337 Neves et al also reported an increase in VEGF expression following irradiation of male Wistar rats
338 with transverse *rectus abdominis* musculocutaneous flap at 660 nm or 830 nm (107). Cheng et al
339 also reported application of 450nm light induced significant increases in COX2 and VEGF in a dose
340 dependent manner (0.001-0.1J/cm²) relative to lipopolysaccharide treated microglial cells (123).
341 Hence, the effects of blue, red and NIR light on NFκB associated pathways have been reported in a
342 handful of studies. However, none to date have evaluated the effects of green light on these
343 pathways. Hence, it will be prudent in the future to evaluate the wavelength and dose dependent
344 effects of light on downstream targets of PBM.

345 3.2 Transforming growth factor-β (TGFβ) signalling

346

347 Transforming growth factor-β (TGF-β) molecules represent a family of growth factors in which
348 there are three mammalian isoforms: TGF-β₁, TGF-β₂ and TGF-β₃. They have been extensively
349 documented for their crucial role in wound healing processes (124) and in promoting angiogenesis
350 and fibrosis (125). They are secreted by a variety of cell types in inactive form as latent-TGF-β in
351 which a TGF-β dimer held together by disulphide bonds is non-covalently bound to a pro-domain
352 known as latency associated peptide (LAP). This complex is also commonly referred to as small latent
353 complex (SLC). The dissociation of this complex to enable activation of free TGF-β can be induced by
354 a range of activation stimuli including heat and pH changes (126, 127). Notably, one mechanism of
355 particular interest here is that PBM could induce activation of TGF-β signalling (128-130). In a recent
356 study Arany et al employed a laser emitting a wavelength of 904 nm with radiant exposure outputs
357 ranging from 0.1-6 J/cm² and concluded that PBM was able to activate latent-TGFβ₁ (131). It has
358 subsequently been hypothesised that light induces an increase in levels of ROS including superoxide
359 (O₂⁻) (132) which interacts with the methionine 253 amino acid residue on LAP (133). This, in turn,
360 then induces a conformational change in LAP, enabling its dissociation from TGF-β enabling it to bind
361 with high affinity to its cell-surface receptors, including TGF-β receptors (TGFβRI, TGFβRII and

362 TGF β RIII). Notably, TGF β RIII binds TGF- β 1 and then transfers it to TGF β RI and TGF β RII, which are
363 both serine/threonine kinases. In turn, these receptors phosphorylate transcription factors including
364 “small mothers against decapentaplegic” (Smad). Once phosphorylated Smad2 and Smad3 bind Smad4,
365 the complex then translocates to the nucleus and interacts with transcriptional coactivators
366 including p300, a nuclear scaffolding protein. This signalling then enables the binding of the complex
367 with the Smad binding element, leading to the transcription of multiple target genes (134).

368 Interestingly, several authors have also provided evidence for an increase in the activity of
369 Smad proteins following irradiation. The Smad family is comprised of the receptor Smads (Smad-1, -
370 2, -3, -5 and -8/-9), the inhibitory Smads (Smad-6 and -7) and the co-Smad, Smad-4. Hirata et al
371 found that irradiation at 805 nm induced increases in phosphorylation of Smad-1/-5/-8 (135).
372 Interestingly, Dang et al also found an increase in phosphorylated Smad proteins, specifically Smad-2
373 and Smad-4 following irradiation at 800 nm (136). Similarly, Yuchao et al reported application of
374 475nm light induced significant increases in Smad2 phosphorylation. Providing evidence that blue
375 light may also show efficacy in modulating TGF β signalling (137). Hence, these data indicate the
376 possible involvement of TGF- β signalling through Smad proteins during the transduction of the
377 molecular effects of PBM. However, other pathways are also induced by TGF- β signalling including
378 the mitogen associated protein kinase pathway (MAPK (138)). Therefore, it will be interesting to
379 determine how PBM, modulates TGF- β signalling through these interlinked pathways, and which
380 wavelengths of light can induce which pathway.

381

382 3.3 Nuclear factor erythroid 2-related factor 2 (Nrf2) signalling

383

384 Nrf2 is a protein in the “basic leucine zipper protein” (bZIP) family and is implicated in regulation of
385 the expression of antioxidant proteins (139). Increases in ROS production lead to the dissociation of
386 Nrf2 from its inhibitor, Keap1, targeting it for degradation. This enables Nrf2 to translocate into the
387 nucleus and induce the transcription of antioxidant genes, due to the binding of Nrf2 to antioxidant
388 response elements (AREs). To date, only a handful of studies have evaluated the effects of PBM on
389 Nrf2 expression and activity. Interestingly, Sohn et al reported an increase in Nrf2 gene expression
390 following irradiation at 635 nm (140). Similarly, Trotter et al also found that application of blue light
391 induced significant increases in Nrf2 expression *in vitro* (141). This acts as a feedback mechanism
392 following NF κ B activation so the interaction of these two pathways may be important in PBM
393 modulation of chronic inflammatory diseases. Indeed a differential upregulation of Nrf2 may be
394 important in such diseases. However, further work will be required to fully dissect the effects of blue
395 and green light on Nrf2 signalling.

396

397 3.4 Mitogen activated protein kinase (MAPK) signalling

398

399 Mitogen activated protein kinases (MAPKs) are a family of serine/threonine protein kinases that play
400 an essential role in the regulation of a diverse number of cellular activities ranging from cell
401 signalling to cell death. There are three subgroups of MAPKs including extracellular signal regulated
402 kinases (ERKs including ERK-1 and ERK-2), p38 MAPKs (p38 α , p38 β , p38 γ and p38 δ) and c-Jun-N-
403 terminal kinases (JNKs including JNK-1, JNK-2 and JNK-3). The three subgroups of MAPKs are finely
404 regulated by a series of different kinases. Their activation is initiated by first the induction of MAPK
405 kinase kinase (MAP3K) which in turn phosphorylates MAPK kinase (MAP2K). This then finally leads to

406 the phosphorylation and activation of the MAPKs. Each MAPK is regulated by specific kinases as
407 described below:

408

409 3.4.1 *Extracellular-regulated kinase (ERK) signalling*

410

411 The activity of the ERK pathway can initially be induced by receptor tyrosine kinases including
412 TGF β R1, a member of the TGF β signalling pathway previously proposed to play a role in transducing
413 the effects of PBM (131). The activation of TGF β R1 enables the downstream activation of Ras-
414 activating protein, which catalyses the phosphorylation of inactive Ras bound to guanosine
415 diphosphate (Ras-GDP) to form active Ras guanosine triphosphate (Ras-GTP). Ras then
416 phosphorylates Raf, which in turn phosphorylates MEK, which ultimately induces phosphorylation of
417 ERK, culminating in gene transcription changes that lead to proliferation, differentiation or
418 apoptosis. Interestingly, several authors have reported the effects of PBM on signalling molecules
419 involved in this pathway. In their study, Kim et al evaluated the response of human outer root
420 sheath cells to PBM at wavelengths of 415 nm, 525 nm, 660 nm or 830 nm (142). Notably they found
421 that PBM induced an increase in ERK phosphorylation.

422

423 3.4.2 *p38 MAPKs*

424

425 The p38 MAPK pathway is activated by an array of stimuli including inflammatory cytokines, heat
426 shock or ligands for G-protein coupled receptors (GPCRs). These stimuli induce the activation of an
427 array of MAP3Ks including TGF β activated kinase-1 (TAK1). Activation of MAP3Ks enables the
428 phosphorylation of MEK3 or MEK6. In turn, these kinases phosphorylate members of the p38 family,
429 inducing their activation and hence a series of downstream effects including modulation of cytokine
430 production and apoptosis (143).

431 Several authors have reported the effects of PBM on p38 MAPK signalling. Interestingly,
432 both Kim et al and Chu et al concluded that red light PBM induced increased p38 phosphorylation
433 and therefore increases in the activity of this pathway (142, 144). However, a further study
434 concluded that following application of red light, there was a decrease in p38 phosphorylation (145).
435 The difference in response may be due to the difference in radiant exposures employed in the
436 different studies. For example, the authors reporting an increase in p38 phosphorylation when using
437 a light source with a radiant exposure output of less than 12 J/cm², whilst, a radiant exposure of 18
438 J/cm² induced a decrease in p38 phosphorylation. Hence, this may show a biphasic dose response in
439 which lower doses of light induce stimulatory effects whilst higher doses cause inhibitory effects.
440 However, further work will be required to evaluate this hypothesis, particularly with reference to
441 blue and green light.

442

443 3.4.3 *c-Jun N-terminal kinase (JNK)*

444

445 The JNK pathway is activated by an array of stimuli including cytokines, growth factors and the
446 ligation of specific receptors. In turn these stimuli activate some of the same MAP3Ks induced in the
447 p38 MAPK pathway including apoptosis signal-regulating kinase 1 (ASK1), mitogen-activated protein

448 kinase kinase kinase (MEKK1) and mitogen-activated protein kinase kinase kinase 3 (MEKK3). In turn,
449 these MAP3Ks can phosphorylate either MKK4 or MKK7, Mitogen-activated protein kinase kinases
450 (MAP2Ks) specific to the JNK pathway. MKK4 and MKK7 can also activate MKK3 and 6, enabling the
451 activation of the p38 MAPK pathway (146).

452 Interestingly, Silva et al explored the effects of 780 nm light at a radiant exposure of 10
453 J/cm² on JNK phosphorylation of mice with diet-induced obesity. They concluded that PBM induced
454 a significant reduction in JNK phosphorylation and hence could prove useful in treating effects
455 induced by a high fat diet (147). Similarly, in a mouse model for depression, Salehpour et al
456 evaluated the effects of 810 nm light at a radiant exposure of 33.3 J/cm² and found that PBM
457 induced reductions in JNK phosphorylation and other members of the MAPK signalling pathway
458 including p38 (148). The authors also reported that treatment induced decreases in the serum levels
459 of key pro-inflammatory cytokines, including Tumour necrosis factor- α (TNF- α). Hence, these data
460 provide evidence that PBM could modulate JNK signalling and therefore downstream effects,
461 including the production of pro-inflammatory cytokines. However, the effects of blue and green light
462 on JNK signalling are yet to be explored, hence future work may endeavour to evaluate the
463 wavelength dependent effects of PBM on the JNK pathway.

464 4 *In vitro and in vivo application of blue and green light PBM therapy*

465

466 4.1 *Introduction*

467

468 Whilst a number of reviews have been published detailing the possible therapeutic efficacy of red
469 and NIR PBM (132, 158), none to date have extensively explored the effects of blue or green light
470 either *in vitro* or *in vivo*. Hence, this section investigates the potential of blue and green light PBM in
471 therapeutic application to determine whether these short wavelengths of light could be efficacious.

472 4.2 *Methods*

473

474 To assess literature surrounding the effects of green and blue light PBM, a systematic review of
475 relevant literature was performed using Scopus. The Scopus database was employed as a means to
476 undergo key word searches to provide an overview of literature regarding the effects of blue and
477 green light PBM. Future work may endeavour to use a broader range of databases including
478 MEDLINE and PubMed to evaluate the effects of blue and green light PBM.

479 The two searches outlined in *Figure 3* were performed separately and a series of key terms
480 and wavelengths were included to refine the search. Following literature searches, the results were
481 filtered for 'articles only' and for articles published within the past ten years (25/1/2008-25/1/2019).
482 This timeline was selected to ensure a manageable sample of articles were included in the review,
483 where key MeSH accepted terms were relevant to articles (including LLLT and PBM) investigated.
484 Key words described in *Figure 3* were input into the Scopus database and systematic evaluation
485 ensured irrelevant articles were excluded from the review. For example, those reporting the use of
486 PBM but using biological assays including 'Alamar blue' (cell metabolic activity assay) or 'trypan blue'
487 (a vital stain used to differentiate between live and dead cells *in vitro*) were not included as they did
488 not specifically report the effects of blue light *in vitro* or *in vivo*. Other exclusion criteria included
489 elimination of articles that reported the effects of lasers on remodelling tissue at a high power. For

490 example, photoselective vaporisation commonly uses lasers emitting green light and the procedure
491 involves the burning away of excess tissue to enable normal urine flow through the prostate (159).
492 As PBM is commonly defined as modulation of tissue response rather than removal of tissue, these
493 articles were excluded. Review articles were also excluded from analysis. Articles selected for review
494 were then assessed in terms of the reporting of light properties (including wavelength, irradiance,
495 radiant exposure, exposure time and beam area), the application of the light source (i.e. *in vitro*, *in*
496 *vivo* or *ex vivo*) and the outcome of each study (therapeutically beneficial or harmful).

497

498

499

500

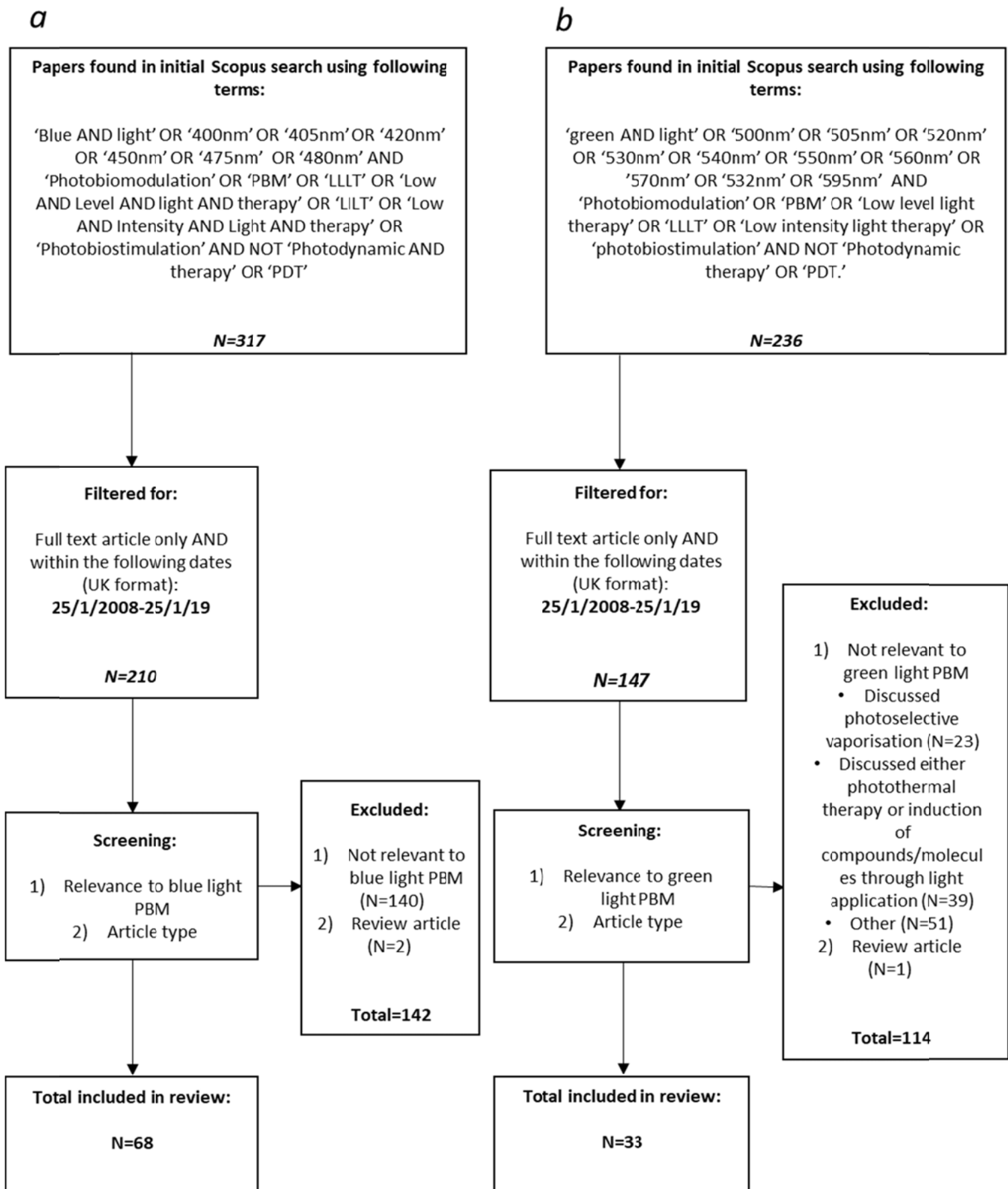


Figure 3: Flow chart describing the strategy employed to identify relevant articles illustrating the effects of a) blue light and b) green light PBM.

502 4.3 Results

503

504 A Scopus search was undergone to identify publications citing key words illustrated in *Figure 3* and
505 published within the following time frame: 25/1/2008-25/1/2019. This would then enable
506 identification of key parameters that may induce beneficial effects *in vivo*.

507 4.3.1 Blue light PBM

508

509 An initial search employing the Scopus database resulted in 317 articles citing the search terms
510 described in *Figure 3a*, articles were then subsequently filtered and screened, and it was concluded
511 that 68 articles were suitable for further review (17-21, 29-31, 64, 68-73, 79, 89, 160-210).

512 Of the articles reviewed, 72% (49/68) reported a positive effect following the application of
513 blue light, with 7% (5/68) reporting negative effects and 21% (14/68) reporting no significant effect.
514 Whilst, the majority of articles within this review reported the effects of PBM on tissue, a handful
515 also evaluated the bactericidal effect of blue light PBM (3/68 (167, 192, 199)). Although, the
516 mechanism of blue light in inducing bacterial cell death is not a focus of this current review, we felt it
517 important to highlight this as a further application of PBM which has therapeutic application (211).
518 Notably however, the parameters required to induce a bactericidal effect ($>55\text{J}/\text{cm}^2$) are much
519 higher than those applied to induce tissue effects ($<55\text{J}/\text{cm}^2$). Hence, when exploring possible
520 beneficial parameters for tissular applications, articles evaluating the antimicrobial properties of
521 blue light were excluded.

522 When evaluating the recording and reporting of treatment parameters, it was found that
523 68% (46/68) of articles failed to report any information regarding light characterisation procedures
524 and relied entirely on the manufacturers reported values. Of the other 22 articles, the light
525 characterisation procedures reported were minimal, where the majority reported either analysis of
526 power or irradiance output of a light source (12/22 (68, 73, 160, 164, 168, 169, 174, 179, 195, 197,
527 204, 206)). Interestingly, these figures are similar to those reported by Hadis et al in which 76% of
528 articles evaluated failed to report the use of any light characterisation techniques (15). A series of
529 other key parameters were also not reported in full by authors including irradiance (29%, 20/68),
530 exposure time (34%, 23/68), radiant exposure (38%, 26/68) and beam area (82%, 56/68).

531 From those articles reporting parameters, median values could be calculated. For example,
532 for those articles reporting a positive effect of PBM, a median value of radiant exposure of $7.8\text{ J}/\text{cm}^2$
533 (range: $1.5\text{--}90\text{ J}/\text{cm}^2$) was determined. In fact, 63% (17/27) of articles reporting both positive effects
534 of PBM used radiant exposure values $<10\text{ J}/\text{cm}^2$. Interestingly, only one article reported a beneficial
535 effect of PBM on tissue at a radiant exposure $>55\text{ J}/\text{cm}^2$ (196). For articles reporting negative effects
536 of PBM, a median radiant exposure of $7.5\text{ J}/\text{cm}^2$ (range: $3\text{--}183.43\text{ J}/\text{cm}^2$) was calculated. However, of
537 the 5 articles reporting a negative effect of PBM only 4 reported radiant exposure values, 3 of which
538 were $>30\text{ J}/\text{cm}^2$. Interestingly of those reporting no significant effect of blue light, an average radiant
539 exposure of $8\text{ J}/\text{cm}^2$ was reported (range: $0.378\text{--}80\text{ J}/\text{cm}^2$). In which, 4/7 of those articles studied
540 reported the radiant exposure utilised in experimentation. Hence, it is apparent from these findings
541 that further work is required to gain a better understanding of the biphasic effect of blue light and
542 also to demonstrate the importance of recording and reporting treatment parameters. *Table 1*
543 summarises the studies evaluated in this review including the study type, reported parameters and
544 outcomes.

545 *Table 1: Citations identified from a review of the literature evaluating the effects of blue light PBM*
546 *using the following search terms: 'Blue AND light' OR '400 nm' OR '405 nm' OR '420 nm' OR '450 nm'*
547 *OR '475 nm' OR '480 nm' AND 'Photobiomodulation' OR 'PBM' OR 'LLLT' OR 'Low AND Level AND*
548 *light AND therapy' OR 'LILT' OR 'Low AND Intensity AND Light AND therapy' OR 'Photobiostimulation'*
549 *AND NOT 'Photodynamic AND therapy' OR 'PDT'*

Citation	Light Source	Dose	Study type	Conclusion
1. Mignon et al, 2018(19)	Source: LED Wavelength (nm): 450, 490, 550, 650 and 850 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 30 (450 nm), 30 (490 nm), 30 (550 nm), 7 (590 nm), 60 (655 nm), 80 (850 nm) Time (s): Energy (J): Radiant exposure (J/cm ²): 0-250 (dependent upon wavelength)	<i>In Vitro</i> : Human reticular and papillary dermal fibroblasts	450nm light at 30J/cm ² induced 50% reductions in cell metabolic activity. 450nm and 500nm induced stronger inhibitory effects on reticular DFs vs papillary DFs. 450nm light induced increases in intracellular ROS production. Blue and NIR light induced changes in some similar gene groups. However, more genes were downregulation following irradiation with blue light compared to NIR. Blue light also downregulated expression of genes associated with the TGF-β pathway.
2.Tani et al, 2018 (191)	Source: LED Wavelength (nm): 405, 635, 808 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 12.59 Time (s): 30 Energy (J): Radiant exposure (J/cm ²): 0.378	<i>In Vitro</i> : human osteoblasts and human mesenchymal stromal cells.	Blue light had no significant effect on molecular signalling. 635nm light could be effective in promoting or improving bone regeneration as shown through molecular analysis.
3.Priglinger et al, 2018 (197)	Source: LED Wavelength (nm): 475, 516, 635 Power (mW): Frequency (Hz): pulsed 2.5 (pulse rate 50%) Spot area (cm ²):	Irradiance (mW/cm ²): 40 Time (s): 600 Energy (J): Radiant exposure (J/cm ²): 24	<i>In Vitro</i> : Stromal Vascular fraction cells	Blue, green and red light did not have a cytotoxic effect on cells. Red and green light induced significant increases in vascular endothelial growth factor expression.
4.Falcone et al, 2018 (175)	Source: LED Wavelength (nm): 453 Power (mW): Frequency (Hz): 5% duty cycle, 100Hz Spot area (cm ²):	Irradiance (mW/cm ²): 10 (cw), 200 (pulsed) Time (s): 1800 Energy (J): Radiant exposure (J/cm ²): 18	<i>In Vivo</i> : effects on inflammation and skin barrier recovery.	Reduced IL-1α following irradiation.
5.Veleska-Stevkoska and Koneski, 2018 (21)	Source: LED Wavelength (nm): 410 and 470 Power (mW): Frequency (Hz): Spot area (cm ²): 1.25	Irradiance (mW/cm ²): 750 Time (s): 10-20 Energy (J): 50-100 Radiant exposure (J/cm ²): 7.5-15	<i>In Vivo</i> : Haemostasis in oral surgery (bleeding from tooth extractions).	Blue light shortens bleeding time from extraction socket.
6. Castellano-Pellicena et al, 2018 (17)	Source: LED Wavelength (nm): 447, 505, 530, 655 and 850 (24 well) or 453, 656 (ex vivo) Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): 1200 (24 well) Energy (J): Radiant exposure (J/cm ²): 453 nm- 2, 656 nm- 30	<i>In vitro and ex vivo</i> : Keratinocytes and human skin epidermis	Blue light stimulated metabolic activity of cultured keratinocytes. Low levels of blue light reduced DNA synthesis and stimulated keratinocyte differentiation. Level of differentiation induced by blue light was reduced in opsin 3 (OPN3) knockdown, suggesting OPN3 may be important in blue light induced restoration of barrier function.
7. Fekrazad et al, 2018(176)	Source: Laser Wavelength (nm): 810, 660, 532, 485, combinations: 810-660, 810-485, 660-532, 660-485 Power (mW): 30-200 (dependent on wavelength) Frequency (Hz): Spot area (cm ²): 0.113-0.18	Irradiance (mW/cm ²): 266 (blue), 266 (green) 167 (red), 1333(NIR) Time (s): 3-24 (dependent upon wavelength) Energy (J): Radiant exposure (J/cm ²): 4 (8 for combination)	<i>In vitro</i> : Mesenchymal stem cells.	Cartilage markers were upregulated by 810nm and 810-485nm light. Red and blue-green irradiation induced expression of COL1. Blue, blue-green and green light irradiation reduced osteocalcin expression. Stimulatory effects on osteogenesis were seen for red and near infra-red lasers but green light had inhibitory effects. Blue light was not reported to induce inhibitory effects. Cons: Parameters differ considerably from one wavelength to the next, particularly when evaluating combination treatments. Making results of which wavelength is most effective questionable.
8. Rocca et al, 2018 (183) (198)	Source: diode laser (450 nm, 635 nm, 808 nm, Er:YAG laser (2940 nm) Wavelength (nm): 2940,	Irradiance (mW/cm ²): 280 (808 nm), 280 (450 nm), 1000 (635 nm) Time (s): 60 (2940 nm),	<i>In Vivo</i> : Human	The diode lasers proved more effective than the Er:YAG in reducing pain scores over a 7 day period. 635nm had the most immediate

	808, 450, 635 Power (mW): 200 (635 nm), 1500 (808 nm), 500 (450nm) Frequency (Hz): 20 (Er:YAG) Spot area (cm²):	3x30 (635 nm), 60 (808 nm), 60 (450 nm) Energy (J): Radiant exposure (J/cm²): 76.43 (Er:YAG), 36 (635 nm), 50 (808 nm), 17 (450 nm)		effect, but there was no significant difference pain score using the 3 diode lasers after 7 days.
9. Wang et al, 2018 (208)	Source: LEDs Wavelength (nm): 405 Power (mW): 200 Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 20 Time (s): 900 Energy (J): Radiant exposure (J/cm²): 12	<i>In Vitro:</i> Whole blood samples (human)	No significant change in the absorption spectra exhibited by blood following irradiation.
10. Kim et al, 2017(187)	Source: LEDs Wavelength (nm): 410, 630, 830 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²):	<i>In Vivo:</i> Mouse model	Wound closure percentage over 10 days was greatest when an 830nm LED used. Increased TGF- β and collagen 1 but downregulated SMAD7.
11. Rohringer et al, 2017 (68)	Source: LED Wavelength (nm): 475, 516, 635 Power (mW): Frequency (Hz): 50% pulse rate, 2.5Hz Spot area (cm²):	Irradiance (mW/cm²): 80 Time (s): 600 Energy (J): Radiant exposure (J/cm²): 24	<i>In Vitro:</i> Human umbilical vein endothelial cells	Red and green light induced proliferation and migration of endothelial cells whilst blue light had no significant impact. Blue light only induced significant increases in ROS production. NOTE: irradiance and irradiation time values do not correlate with fluency. Cells irradiated at 20% confluency – may explain negative effects of blue light.
12. Wang et al, 2017 (64)	Source: LED array (415 nm), Filtered lamp (540 nm), Diode laser (660 nm and 810 nm) Wavelength (nm): 415, 540, 660, 810 Power (mW): Frequency (Hz): Spot area (cm²): 4	Irradiance (mW/cm²): 16 Time (s): 188 Energy (J): Radiant exposure (J/cm²): 3	<i>In vitro:</i> human adipose-derived stem cells.	Blue and green light induce significant increases in intracellular calcium and ROS, reduce mitochondrial membrane potential, lower intracellular pH and reducing cellular proliferation. Red and NIR light have the opposite effect. Labelled fluency as irradiance. Different delivery systems may alter light delivery (coherent vs non-coherent light sources).
13. Wang et al, 2017(69)	Source: LED Wavelength (nm): 480, 560, 660 and white (400-750) Power (mW): 3000 Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 250 Time (s): Energy (J): Radiant exposure (J/cm²):	<i>In vitro and in vivo:</i> Irradiated fertile broiler eggs and isolated skeletal muscle and satellite cells.	Green PBM promoted muscle growth and satellite cell proliferation through insulin growth factor-1 signalling in late embryogenesis.
14. Choe et al, 2017 (171)	Source: LED Wavelength (nm): 622, 535, 462 Power (mW): 24000 Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 16 Time (s): 600-1800 (daily) Energy (J): Radiant exposure (J/cm²):	<i>In Vitro:</i> HeLa cells (cancer cell line).	Blue light and high frequency ultrasound induced significant reductions in cell density when compared to red and green light combined with ultrasound. This could be beneficial in alleviating cancer cell proliferation.
15. Buscone et al, 2017 (170)	Source: LED Wavelength (nm): 453, 689 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²): 3.2	<i>In vitro and ex vivo:</i> hair growth and outer root sheath cells.	Blue light at low radiant exposure stimulate hair growth ex vivo.
16. Santos et al, 2017 (70)	Source: LED Wavelength (nm): 405 Power (mW): Frequency (Hz): Spot area (cm²): 0.27	Irradiance (mW/cm²): 300 Time (s): 30-60 Energy (J): Radiant exposure (J/cm²):	<i>In vitro:</i> Subventricular zone (SVZ) cell culture.	Blue light induced transient increases in ROS, causing increased neuronal differentiation and increases retinoic acid receptor levels. The effects are heightened with the addition of light reactive nanoparticles.
17. Fekrazad et al, 2017(18)	Source: laser: GaAs (405 nm, 532 nm), InGaAlP (660 nm) and GaAlAs (810 nm) Wavelength (nm): 405, 532, 660, 810 Power (mW): Frequency (Hz): Spot area (cm²): 1	Irradiance (mW/cm²): 200 Time (s): Energy (J): Radiant exposure (J/cm²): 1.5	<i>In Vivo:</i> Male Wistar rats (n=60)	Green, blue, red and infrared light irradiation may accelerate healing process.
18. Lee et al, 2017(190)	Source: LED Wavelength (nm): 410, 630, 830 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 205 (405nm), 172 (630), 50 (830) Time (s): Energy (J): Radiant exposure (J/cm²):	<i>In Vitro:</i> Keloid fibroblasts	Blue did not affect cell viability. COL1 gene and protein expression decreased significantly after irradiation with blue light and may be effective in preventing keloid formation.

		10		
19. Alba et al, 2017 (29)	Source: LED (470) and laser (660) Wavelength (nm): 470 and 660 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): 180 (470), 60 (660) Energy (J): 6-8 Radiant exposure (J/cm ²):	<i>In Vivo</i> : treatment of acne vulgaris	The combined use of red and blue light proved beneficial in reducing inflammation and enhancing wound healing when compared to the use of salicylic acid for treatment.
20. Mignon et al, 2017(193)	Source: LED Wavelength (nm): 400, 500, 530, 590, 655, 850 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 3-80 Time (s): Energy (J): Radiant exposure (J/cm ²): 2-30	<i>In Vitro</i> : Human dermal fibroblasts	The effects of blue light on cell metabolism were dramatically influenced by FBS concentration, confluency level of cells and the fluency values applied to cells.
21. Yoshimoto et al, 2017(210)	Source: LED Wavelength (nm): 465 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 30 Time (s): 1800 Energy (J): Radiant exposure (J/cm ²): 54	<i>In Vitro</i> : Human colon cancer cells (HT-29 or HCT-116)	Blue light irradiation reduced cancer cell viability. However, this effect was reversed in an Opsin 3 (Opn3) knockdown.
22. Yuan et al, 2017(71)	Source: LED Wavelength (nm): 470 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 20 Time (s): 60-3600 Energy (J): Radiant exposure (J/cm ²):	<i>In Vitro</i> : Bone marrow-derived mesenchymal stem cells (BMSCs)	Blue light inhibited osteogenic differentiation, induced apoptosis as a result of increased ROS production and DNA damage.
23. Monrazeri et al, 2017 (194)	Source: LED Wavelength (nm): 630, 808, 450 Power (mW):100 (630 nm and 808 nm), 3000 (450 nm) Frequency (Hz): Spot area (cm ²): 1	Irradiance (mW/cm ²): 100 (630 nm) Time (s): Energy (J): 48J per point Radiant exposure (J/cm ²):	<i>In vivo</i> : Human	Combining all three wavelengths reduced abdominal girth significantly.
24. Li et al, 2016(191)	Source: LED Wavelength (nm): 630, 460 Power (mW):100 (630nm and 808nm), 3000 (450nm) Frequency (Hz): Spot area (cm ²): 300	Irradiance (mW/cm ²): 50 Time (s): 900-1800 Energy (J): Radiant exposure (J/cm ²): 45-90	<i>In vivo</i> : Japanese big ear white rabbits, induced wound model (incisions in back)	Red light was more effective in promoting wound healing than blue light.
25. Khori et al, 2016 (186)	Source: Laser Wavelength (nm): 405, 532, 632 Power (mW): 1-3 Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): 600 (10 treatments, 3 times a week) Energy (J): Radiant exposure (J/cm ²):	<i>In vivo and in vitro</i> : BALB/c inbred female mice and mouse mammary carcinoma cell line (4T1).	Blue light reduced tumour volume and gene expression markers for tumorigenesis.
26. AlGhamdi et al, 2016 (161)	Source: laser Wavelength (nm): 457, 635, 355 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 25 Time (s): 80 Energy (J): Radiant exposure (J/cm ²): 2	<i>In Vitro</i> : Melanocytes from normal human melanocytes.	PBM at all wavelengths induced the production of stage I melanosomes to the highest levels relative to control cells. In particular, red and blue laser PBM induced the highest increase in % level of stage I melanosomes. This indicates significant stimulation of melanogenesis.
27. Wang et al, 2016 (209)	Source: LED array (420), Filtered lamp (540), Diode laser (660, 810). Wavelength (nm): 420, 540, 660, 810 Power (mW): Frequency (Hz): Spot area (cm ²): 4	Irradiance (mW/cm ²): 16 Time (s): 188 (five times, every 2 days). Energy (J): Radiant exposure (J/cm ²): 3	<i>In Vitro</i> : Human adipose-derived stem cells.	Blue and green light were effective in stimulating osteoblast differentiation and increasing intracellular calcium levels than red and near infra-red light. Blue and green light could activate light-gated calcium ion channels.
28. Masson-Meyers et al, 2016(30)	Source: LED Wavelength (nm): 470 Power (mW): 150 Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 30 Time (s): Energy (J): Radiant exposure (J/cm ²): 3, 5, 10, 55	<i>In Vitro</i> : Human Dermal Fibroblasts	Blue light and radiant exposure of 5J/cm ² improved wound healing, increased protein concentration and reduced IL-6 secretion significantly. There was no effect of irradiation on cell viability.
29. Ashworth et al, 2016 (164)	Source: LED Wavelength (nm): 450, 510, 660, 860 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): Energy (J): Radiant exposure (J/cm ²): Other: photons/cm ² /s	<i>Ex Vivo</i> : Rat or mouse spinal cord slices	All four wavelengths at the highest intensity output reduced immunoreactivity.
30. Figurova et al, 2016 (178)	Source: LED Wavelength (nm): 685/470 combined Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 8 Time (s): Energy (J): Radiant exposure (J/cm ²): 3.36	<i>In Vivo</i> : Minipigs	Combined red and blue light therapy induced improved tissue healing relative to control groups.

31. Becker et al, 2016 (165)	Source: LED Wavelength (nm): 453 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 23 Time (s): 1800 Energy (J): Radiant exposure (J/cm ²):	<i>In Vitro</i> : Melanoma cells	The effects of blue light on cell viability were dose dependent and blue light down regulated anti-inflammatory genes but upregulated genes associated with apoptosis. Significant decreases in viability were witnessed after irradiation times of 1800s.
32. Dereci et al, 2016 (173)	Source: LED (blue, 400-490), GaAlAs diode laser (NIR, 980nm) Wavelength (nm): 400-490, 980 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 12 (400-490), 200 (980) Time (s): Energy (J): Radiant exposure (J/cm ²): 13 (400-490, 20 (980)	<i>In Vivo</i> : Wistar rats	Whilst high doses of blue light were inhibitory, low doses proved efficacious in promoting bone regeneration to similar levels to NIR light.
33. Takhtfooladi and Sharifi, 2015 (204)	Source: GaAlAs (680), LED (650, 450) Wavelength (nm): 680, 650, 450 Power (mW): 10 (680) Frequency (Hz): Pulsed (no info, 680 only) Spot area (cm ²): 0.4 (680), 1.5 (650, 450)	Irradiance (mW/cm ²): Time (s): 200s (680), 600s (450, 650) 14 days Energy (J): Radiant exposure (J/cm ²): 10 (680), 650 (2.4), 450 (2.4)	<i>In Vivo</i> : New Zealand rabbits	Blue and red LEDs had no significant effect on cell proliferation or myelination. Conversely, laser red light had a significant effect. This may be due to the pulsed modality of the laser light source.
34. Fekrazad et al, 2015 (177)	Source: laser Wavelength (nm): 630, 532, 425 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 50 (630 nm and 532 nm), 55 (425 nm) Time (s): Energy (J): Radiant exposure (J/cm ²): 2	<i>In Vivo</i> : Diabetes induced male Wistar rats	All three wavelengths induced significant increases in wound healing, where red light was most effective.
35. Masson-Meyers et al, 2015 (192)	Source: LED or Laser Wavelength (nm): 405 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): 900, 1800, 14400 Energy (J): Radiant exposure (J/cm ²): 40, 54, 81, 121	<i>In Vitro</i> : Methicillin Resistant Staphylococcus Aureus (MRSA)	Both LED and laser proved efficacious in suppressing bacterial growth to significant levels at all four radiant exposure values evaluated.
36. Schafer and McNeely, 2015 (199)	Source: LED Wavelength (nm): 405 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 30 Time (s): Energy (J): Radiant exposure (J/cm ²):	<i>In Vitro</i> : Staphylococcus Epidermis, Staphylococcus Aureus and Propionibacterium Acnes	The effects of blue light combined with ultrasound were dose dependent where it is proposed that bacterial cells become more susceptible to the antimicrobial effects of blue light following ultrasound application.
37. Niu et al, 2015 (195)	Source: LED Wavelength (nm): 405, 630, 660 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 161μW/cm ² nm (405 nm), 300μW/cm ² nm (630 nm), 545μW/cm ² nm (660 nm) Time (s): 600 Energy (J): Radiant exposure (J/cm ²): 1.604 (405 nm), 3.409 (630 nm), 6.538 (660 nm)	<i>In Vitro</i> : Keratinocytes	The combination of blue light, red light and curcumin was able to regulate proliferation and apoptosis of keratinocytes. Without curcumin, light did not influence cell viability.
38. AlGhamdi et al, 2015 (162)	Source: diode laser Wavelength (nm): 355, 457, 635 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 25 Time (s): 20-200 Energy (J): Radiant exposure (J/cm ²): 0.5-5	<i>In Vitro</i> : Melanocytes	Blue laser proved most efficacious in promoting cell proliferation and migration.
39. Pfaff et al, 2015 (196)	Source: LED Wavelength (nm): 453 Power (mW): Frequency (Hz): High (200mW/cm ²) and low (100mW/cm ²) duty cycles employed Spot area (cm ²):	Irradiance (mW/cm ²): 100 (low), 200 (high) Time (s): 1800 Energy (J): Radiant exposure (J/cm ²): 90	<i>In Vivo</i> : Treatment of patients with mild Psoriasis Vulgaris (Pv).	Blue light proved to significantly reduce Pv severity at both irradiance outputs.
40. Bumah et al, 2015(167)	Source: LED Wavelength (nm): 470 Power (mW): 150 (18 delivered to cultures) Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 30 Time (s): Energy (J): Radiant exposure (J/cm ²): 55	<i>In Vitro</i> : MRSA	Blue light alone is effective in suppressing MRSA growth, where there was no significant difference in the effect of blue light and the combination of blue light and hyperbaric oxygen.
41. Jung et al, 2015 (183)	Source: LED Wavelength (nm): 415, 630 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): 76-615 Energy (J): Radiant exposure (J/cm ²): 5-40	<i>In Vitro</i> : Human Sebocytes	Blue and red light influence lipid production and may have beneficial effects on acne through the suppression of sebum production.
42. Teuschl et al, 2015 (206)	Source: LED Wavelength (nm): 470, 630	Irradiance (mW/cm ²): 50 Time (s): 600 (5 times, once per day)	<i>In Vitro</i> : C2C12 (myoblast), NIH/3T3 (fibroblast), BICR10	Blue light reduced cell proliferation and promoted necrosis. Red light promoted cell proliferation and

	Power (mW): 1000 Frequency (Hz): Spot area (cm²):	Energy (J): Radiant exposure (J/cm²): 30	(keratinocytes)	increased rate of wound healing.
43. Hadis et al, 2015(181)	Source: LED Wavelength (nm): 400-900 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 3.5 Time (s): 15-120 Energy (J): Radiant exposure (J/cm²): 0.05-0.42	<i>In Vitro:</i> Dental Pulp cells (DPCs)	Blue light had no significant effect on DPCs whilst wavelengths of 625nm, 660nm, 789nm and 800nm induced significant increases in mitochondrial activity. Particularly after 24hrs and irradiation periods of 30s.
44. De Sousa et al, 2015 (31)	Source: LED Wavelength (nm): 450 Power (mW): 70 Frequency (Hz): Spot area (cm²): 0.00785	Irradiance (mW/cm²): Time (s): 0-343 Energy (J): Radiant exposure (J/cm²): 3-24	<i>In Vitro:</i> <i>Staphylococcus aureus</i> , <i>Pseudomonas aeruginosa</i>	Blue light inhibited bacterial growth at fluency values greater than 6J/cm ² .
45. Gold et al, 2014 (180)	Source: LEDs Wavelength (nm): 405-460 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²):	<i>In Vivo:</i> Human	Induced a reduction in acne vulgaris inflammatory lesions. Did induce increases in skin temperature up to 41°C.
46. Schoenly et al, 2014(200)	Source: laser Wavelength (nm): 400 Power (mW): Frequency (Hz): 60ns laser pulse Spot area (cm²):	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²): <8	<i>In Vitro:</i> Human teeth	Removal of calculus is thickness dependent and can occur at radiant exposure <5J/cm ²
47. Buravlev et al, 2014 (73)	Source: LED Wavelength (nm): 442 Power (mW): 70 Frequency (Hz): Spot area (cm²): 0.00785	Irradiance (mW/cm²): Time (s): 30-300 Energy (J): Radiant exposure (J/cm²): 30	<i>In Vitro:</i> Mitochondria isolated from male albino rat livers.	Blue light restored nitric oxide inhibited rates of respiration to normal. It is hypothesised blue light irradiation induces photolytic destruction of nitrosyl complexes that inhibit the activities of complex I and III of the electron transport chain.
48. Sinclair et al, 2014 (203)	Source: LED Wavelength (nm): 465, 574 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 0.0848 (blue), 0.0185 (yellow) Time (s): 2700 Energy (J): Radiant exposure (J/cm²): Other: 68 lux, 1.21xphotons/cm ² /s	<i>In Vivo:</i> Patients with traumatic brain injury (TBI)	Blue light is effective in alleviating fatigue and daytime sleeping following TBI
49. Hochman et al, 2014 (182)	Source: LED (470 nm and 660 nm) and Laser (660 nm and 808 nm, no details of laser source) Wavelength (nm): 470, 660, 808 Power (mW): 100 (808nm, 660nm Laser) and 350 (470nm and 660nm LED). Frequency (Hz): Spot area (cm²): 0.5 (LED), 0.028 (laser)	Irradiance (mW/cm²): Time (s): 114 (LED), 396 (laser) Energy (J): 40 (both) Radiant exposure (J/cm²): 80 (LED), 1429 (laser)	<i>In Vivo:</i> Skin of adult male Wistar rats.	Infrared (808nm) laser irradiation enhances neuropeptide secretion in healthy rat skin, whilst other sources of light and wavelengths had no significant impact.
50. Dungal et al, 2014(174)	Source: LED Wavelength (nm): 470, 629 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 50 Time (s): 600 Energy (J): Radiant exposure (J/cm²): 30	<i>In Vivo:</i> Sprague-Dawley rats	Both wavelengths promoted angiogenesis, improved tissue perfusion, reduced tissue necrosis and therefore promoted wound healing.
51. KazemiKhood and Ansari et al, 2014 (185)	Source: Optical fiber Wavelength (nm): 405, 632.8 Power (mW): 1.5 Frequency (Hz): Spot area (cm²): 0.01	Irradiance (mW/cm²): Time (s): 1800 (every other day, 14 sessions) Energy (J): Radiant exposure (J/cm²):	<i>In Vivo:</i> Intravascular laser irradiation of blood in type 2 diabetic patients and measurements of changes in blood sugar.	Both wavelengths induced significant decreases in blood sugar levels.
52. Burvalev et al, 2014 (168)	Source: Laser (442 nm, 532 nm) and LED (650 nm) Wavelength (nm): 442, 532, 650 Power (mW): 20 Frequency (Hz): Spot area (cm²): 1.57	Irradiance (mW/cm²): 30 Time (s): 30-300 Energy (J): Radiant exposure (J/cm²): 3-31	<i>In Vitro:</i> Mitochondria isolated from rat liver.	Laser of mitochondria at 442nm restored mitochondrial respiration inhibited by NO. Blue light also restored complex IV activity but not complexes I-III. Other wavelengths had no significant effect.
53. Turrioni et al, 2013(207)	Source: Wavelength (nm): 450, 630, 850 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²):	<i>In Vitro:</i> Human dentin	All three wavelengths passed through the dentin barrier. LED power loss and transmittance varied dependent upon dentin thickness and wavelength.

54. Kazemi Khoo et al, 2013(184)	Source: laser Wavelength (nm): 405 Power (mW): 1.5 Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): 1800 Energy (J): Radiant exposure (J/cm ²):	<i>In Vivo</i> : Human diabetic patients	Resulted in modulation of metabolites associated with type 2 diabetes following intravenous PBM.
55. Cheon et al, 2013 (79)	Source: LED Wavelength (nm): 470, 525, 633 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 3.55 (470nm), 4.02 (525nm) and 6.78 (633nm) Time (s): 3600 (9 days) Energy (J): Radiant exposure (J/cm ²):	<i>In Vivo</i> : Sprague Dawley rats and histological analysis	Blue and green light promoted wound healing significantly. Red light promoted collagen production.
56. Burvalev et al, 2013 (169)	Source: HeCd laser (442 nm) diode pumped solid state laser (532 nm) and LED (650 nm) Wavelength (nm): 442, 532, 650 Power (mW): 20 Frequency (Hz): Spot area (cm ²): 1.57	Irradiance (mW/cm ²): Time (s): 30-300 (1 treatment) Energy (J): Radiant exposure (J/cm ²):	<i>In Vivo</i> : Lipopolysaccharide B was applied through intraperitoneal injection to outbred albino rats. Mitochondria were then isolated from rat liver.	Blue light induced a 40% increase in mitochondrial respiration from LPS treated animals at a dose of 6J/cm ²
57. Kushibiki et al, 2013 (72)	Source: Laser Wavelength (nm): 405, 664, 808 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 100 Time (s): 60-120 Energy (J): Radiant exposure (J/cm ²):	<i>In Vitro</i> : mouse preadipocytes (3T3-L1), prechondrocytes (ATDC5), myoblasts (C2C12), mesenchymal stromal cells (KUSA-A1), lung cancer cells (LLC), insulinoma cells (MIN6), fibroblasts (NIH-3T3), human cervix adenocarcinoma cells (HeLa), macrophages differentiated from lymphocytes (THP-1) after treatment with phorbol ester, and rat basophilic leukemia cells (RBL-2H3)	After blue light irradiation, intracellular ROS production was significantly increased in all cell types whilst red and near infra-red light had no significant effect.
58. Fushimi et al, 2012(179)	Source: LED Wavelength (nm): 456, 518, 638 Power (mW): 7560 (638nm), 6930 (456 nm) and 6840 (518 nm) Frequency (Hz): Spot area (cm ²): 30	Irradiance (mW/cm ²): 0.75 (638 nm), 0.25 (456 nm) and 0.17 (518 nm) Time (s): 1200 Energy (J): Radiant exposure (J/cm ²): 0.6 (638 nm), 0.3 (456 nm), 0.2 (518 nm)	<i>In Vivo and in vitro</i> : Induced wound model in ob/ob mice	LED irradiation induced significant increases in growth factor and cytokine secretion. Green LEDs promote wound healing by inducing migratory and proliferative mediators.
59. Lavi et al, 2012(189)	Source: LED Wavelength (nm): 400-505, 600-800 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 30 (600-800 nm), 10 (400-505 nm) Time (s): Energy (J): Radiant exposure (J/cm ²):	<i>In Vitro</i> : Sperm membranes	Visible (especially blue) light induce increase in ROS production in isolate sperm isolated plasma membranes
60. Adamskaya N et al, 2011(160)	Source: LED Wavelength (nm): 470, 630 Power (mW): 1000 Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 50 Time (s): 600 Energy (J): Radiant exposure (J/cm ²):	<i>In Vivo</i> : Induced wound model (excision wound on dorsum), Sprague Dawley rats	Blue light was effective in inducing wound healing and promoting keratin expression.
61. Shuvaeva et al, 2011(202)	Source: laser Wavelength (nm): 473, 650 Power (mW): 20 Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 20 Time (s): Energy (J): Radiant exposure (J/cm ²):	<i>In vivo</i> : WKY and SHR rats	Irradiation with red light proved more effective than blue light in augmenting the constrictive effects of Norepinephrine on pial arteries. However both exerted a significant difference relative to the control.
62. Bonatti et al, 2011(166)	Source: LED Wavelength (nm): 470 Power (mW): 100 Frequency (Hz): Spot area (cm ²): 0.8	Irradiance (mW/cm ²): 125 Time (s): 60-180 Energy (J): 6, 12, 18 Radiant exposure (J/cm ²): 59.87, 122.3, 183.43	<i>In vitro</i> : Keloid and skin fibroblasts, human	Reduced skin fibroblasts following irradiation at 183.43J/cm ² but induced no significant effect on keloid fibroblast number.
63. Ankri et al, 2010(89)	Source: LED Wavelength (nm): 400-830 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): Energy (J): Radiant exposure (J/cm ²):	Computational model of human dermis: photon migration model	480nm may be useful for treating infected wounds whilst 780nm has a higher penetration depth and therefore may be useful for wound healing.
64. De Sousa et al, 2010(172)	Source: LED Wavelength (nm): 700,	Irradiance (mW/cm ²): 7.46 (700 nm), 3.98 (530 nm),	<i>In Vivo and In vitro</i> : Male Wistar rats with	Green and red LEDs induced increases in fibroblast number relative to the

	530, 460 Power (mW): 15 (700nm), 8 (530nm), 22 (460nm) Frequency (Hz): Spot area (cm²): 2.01	10.94 (460 nm) Time (s): 668 (700 nm), 1250 (530 nm), 456 (460 nm) Energy (J): Radiant exposure (J/cm²): 10	excisional wound, followed by histological analysis.	control.
65. Ankri et al, 2010(163)	Source: LED Wavelength (nm): 400- 800 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 130 Time (s): 300 Energy (J): Radiant exposure (J/cm²):	<i>In vitro:</i> Sperm and endothelial cells	Illumination induced increase in NO concentration, particularly blue light.
66. Kushibiki et al, 2010(188)	Source: LED Wavelength (nm): 405 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 100 Time (s): 180 Energy (J): Radiant exposure (J/cm²):	<i>In vitro:</i> Prechondrogenic cells	Intracellular ROS increased and mRNA levels relating to chondrogenesis were elevated.
67. Sebbe et al, 2009(201)	Source: LED Wavelength (nm): 472 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 1.26- 4.73 Time (s): 8-24h Energy (J): Radiant exposure (J/cm²):	<i>In vivo:</i> Male Wistar rats.	Increased bilirubin degradation, important for neonatal jaundice.
68. Tamarova et al, 2009 (205)	Source: LED Wavelength (nm): 480- 3400 (range of source, evaluated 'red, orange, yellow, blue, green, violet') Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 40 Time (s): 600 Energy (J): Radiant exposure (J/cm²): 2.4 (per minute)	<i>In vivo:</i> Male albino rats with area of pain induced by saline injection	Red light was more effective in inducing an analgesic effect. However, all colours induced significant increases in analgesia relative to control.

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567 4.3.2 Green light PBM

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569 A second Scopus database search was undertaken to evaluate the effects of green light PBM using
570 the terms described in *Figure 3b*. An initial search resulted in 236 articles being identified and these
571 articles were subsequently screened for suitability, which then identified 32 relevant articles for
572 further review (18, 32, 33, 64, 68, 69, 79, 164, 168, 171, 176, 177, 186, 197, 205, 209, 212-227).

573 When evaluating the outcomes of studies reporting the effects of PBM it was found that
574 75% (24/32) reported a beneficial effect of green light, whilst 9% (3/32) reported negative effects
575 and 16% (5/32) reported no significant response. Interestingly, this review also included an article
576 evaluating the effect of green light PBM on microbial cell death (219) with high radiant exposures
577 also being used ($\leq 172.8 \text{ J/cm}^2$). These findings further support the use of other visible wavelengths
578 of light in applications other than modulation of tissue response. As described previously this article
579 was also excluded from evaluation of parameters for suitable application to tissue PBM.

580 Exploration of treatment parameters revealed that 72% (23/32) articles failed to report any
581 characterisation protocols or relied entirely upon the parameters stated by the manufacturer. In
582 fact, only one article reported the use of beam profiling to accurately calculate beam area and to
583 provide representative images of the distribution of spectral irradiance (214). Similar to reports in
584 section 4.3.1, a series of key parameters were also not always reported including irradiance (41%,
585 13/32), radiant exposure (44%, 14/32) and beam area (66%, 21/32). In the articles reporting
586 treatment parameters, median treatment values were also determined. In the articles reporting a
587 positive effect of green light PBM, there was a median radiant exposure output of 4 J/cm^2 (range:
588 $0.00362\text{-}30 \text{ J/cm}^2$). Interestingly, it was also found that the most commonly employed wavelength
589 used in these studies was 532 nm, and 35% (11/32) of studies reported the use of this wavelength.
590 Further information detailing parameters and study types employed by authors reviewing the effects
591 of green light PBM are provided in *Table 2*.

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606 Table 2: Citations identified from a review of the literature evaluating the effects of green and yellow
607 light PBM using the following search terms: 'green AND light' OR '500 nm' OR '505 nm' OR '520 nm'
608 OR '530 nm' OR '540 nm' OR '550 nm' OR '560 nm' OR '570 nm' OR '532 nm' OR '595 nm' AND
609 'Photobiomodulation' OR 'PBM' OR 'Low level light therapy' OR 'LLLT' OR 'Low intensity light therapy'
610 OR 'photobiostimulation' AND NOT 'Photodynamic therapy' OR 'PDT.'

Citation	Light Source	Dose	Study type	Conclusion
1. Priglinger et al, 2018 (197)	Source: LED cluster lamp Wavelength (nm): 475, 516, 635 Power (mW): Frequency (Hz): 2.5 (pulsed, 50% rate) Spot area (cm ²):	Irradiance (mW/cm ²): 40 Time (s): 120-300 Energy (J): Radiant exposure (J/cm ²): 24	In Vitro: Adipose tissue derived stromal vascular fraction cells.	Green and red light resulted in increased vascular tube formation and increased concentration vascular endothelial growth factor (VEGF) concentration. Blue light had no significant effect.
2. Askhadulin et al, 2018 (213)	Source: Laser Wavelength (nm): 365, 525, 635 Power (mW): 1-2 Frequency (Hz): 80 (635 nm) Spot area (cm ²): 8 (635 nm)	Irradiance (mW/cm ²): 5000 (635 nm) Time (s): 120 (365 nm, 635 nm) 300 (525nm) , 6 sessions of each Energy (J): Radiant exposure (J/cm ²):	In Vivo: Human	Reduced ulcer healing time and adapted physiological responses ultimately preventing relapse.
3. Fekrazad et al, 2018 (176)	Source: Laser Wavelength (nm): 810, 660, 532, 485, combinations: 810-660, 810-485, 660-532, 660-485 Power (mW): 30-200 (dependent on wavelength) Frequency (Hz): Spot area (cm ²): 0.113-0.18	Irradiance (mW/cm ²): 266 (blue), 266 (green) 167 (red), 1333(NIR) Time (s): 3-24 (dependent upon wavelength) Energy (J): Radiant exposure (J/cm ²): 4 (8 for combination)	In vitro: Mesenchymal stem cells.	Cartilage markers were upregulated by 810nm and 810-485nm light. Red and blue-green irradiation induced expression of COL1. Blue, blue-green and green light irradiation reduced osteocalcin expression. Stimulatory effects on osteogenesis were seen for red and near infra-red lasers but green light had inhibitory effects. Blue light was not reported to induce inhibitory effects.
4. Oh et al, 2018 (225)	Source: LED Wavelength (nm): 630, 595, 480, 410 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 5.47 (410 nm), 13.2 (480 nm), 5.8 (595 nm), 8.63 (630 nm) Time (s): 410nm: 181.2-1816.8 480nm:69.6-742.2 595nm:151.2-1706.4 630nm:93-1146 Energy (J): Radiant exposure (J/cm ²): 1-10	In Vitro: Human umbilical vein endothelial cells (HUVEC)	Irradiation at 630nm induced increases in cell proliferation, NO secretion and eNOS expression from HUVECs. Only evaluated effects on proliferation using other wavelengths where no significant change was witnessed.
5. Rohringer et al, 2017 (68)	Source: LED Wavelength (nm): 475, 516, 635 Power (mW): Frequency (Hz): 50% pulse rate, 2.5Hz Spot area (cm ²):	Irradiance (mW/cm ²): 80 Time (s): 600 Energy (J): Radiant exposure (J/cm ²): 24	In Vitro: Human umbilical vein endothelial cells	Red and green light induced proliferation and migration of endothelial cells whilst blue light had no significant impact. Blue light only induced significant increases in ROS production.
6. Wang et al, 2017(64)	Source: LED array (415 nm), Filtered lamp (540 nm), Diode laser (660 nm and 810 nm) Wavelength (nm): 415, 540, 660, 810 Power (mW): Frequency (Hz): Spot area (cm ²): 4	Irradiance (mW/cm ²): 16 Time (s): 188 Energy (J): Radiant exposure (J/cm ²): 3	In vitro: human adipose-derived stem cells.	Blue and green light induce significant increases in intracellular calcium and ROS, reduce mitochondrial membrane potential, lower intracellular pH and reducing cellular proliferation. Red and NIR light have the opposite effect. Blue and green light inhibit proliferation through activation of TRPV1.
7. Baek et al, 2017 (214)	Source: laser Wavelength (nm): 532 Power (mW): 300 Frequency (Hz): 0.2 (1s) Spot area (cm ²): diameter 1mm	Irradiance (mW/cm ²): Time (s): 10-300 Energy (J): Radiant exposure (J/cm ²):	In vitro: vascular smooth muscle cells	Inhibited platelet derived growth factor-BB induced proliferation and migration. Also induced apoptosis via the p38 MAPK pathway.
8. Wang et al, 2017 (69)	Source: LED Wavelength (nm): 480, 560, 660 and white (400-750) Power (mW): 3000 Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 250 Time (s): Energy (J): Radiant exposure (J/cm ²):	In vitro and in vivo: fertile broiler eggs were irradiated and satellite cells were isolated.	Green light promoted muscle growth and satellite cell proliferation which may be due to an increase in signalling through the insulin growth factor (IGF-1) pathway.

9. Choe et al, 2017 (171)	Source: LED Wavelength (nm): 622, 535, 462 Power (mW): 24000 Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 16 Time (s): 600-1800 (daily) Energy (J): Radiant exposure (J/cm²):	<i>In Vitro:</i> HeLa cells (cancer cell line).	Blue light and high frequency ultrasound induced significant reductions in cell density when compared to red and green light combined with ultrasound. This could be beneficial in alleviating cancer cell proliferation. Green light also drove decreases in cell density but not significantly.
10. Fekrazad et al, 2017(18).	Source: laser: GaAs (405 nm, 532 nm), InGaAlP (660 nm) and GaAlAs (810 nm) Wavelength (nm): 405, 532, 660, 810 Power (mW): Frequency (Hz): Spot area (cm²): 1	Irradiance (mW/cm²): 200 Time (s): Energy (J): Radiant exposure (J/cm²): 1.5	<i>In Vivo:</i> Male Wistar rats (n=60)	Green, blue, red and infrared light irradiation may accelerate healing process.
11. Moskvina et al, 2017 (222)	Source: LAMSIK® device, external pulsed laser (635), intravenous laser blood illumination (ILBI, 365-405 nm and 520-525 nm) Wavelength (nm): 635, 365-405, 520-525 Power (mW): 40000 (635 nm) Frequency (Hz): pulsed 635nm Spot area (cm²): 8 (635 nm)	Irradiance (mW/cm²): Time (s): 12 sessions, 120s (per point, 635 nm), 120s (365-405 nm), 300s (520-525 nm) 6 sessions each alternate. Energy (J): Radiant exposure (J/cm²):	<i>In vivo:</i> Treatment of patients with chronic venous diseases	Reduced time for wound cleansing, stimulates proliferation and epithelialisation processes.
12. Khorri et al, 2016 (186)	Source: Laser Wavelength (nm): 405, 532, 632 Power (mW): 1-3 Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): 600 (10 treatments, 3 times a week) Energy (J): Radiant exposure (J/cm²):	<i>In vivo and in vitro:</i> BALB/c inbred female mice and mouse mammary carcinoma cell line (4T1).	Blue light reduced tumour volume and gene expression markers for tumorigenesis.
13. Roche et al, 2017 (226)	Source: laser diodes Wavelength (nm): 532 Power (mW): 17 (per diode, 170 total) Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): 0.03 Time (s): 1800 (3 times weekly) Energy (J): Radiant exposure (J/cm²): 0.03 (per treatment), 0.36 (in total) calculations appear wrong.	<i>In Vivo:</i> Obese but otherwise healthy individuals, RCT	Reduced circumference of hips, waist and upper abdomen when applied to individuals with a body mass index (BMI) between 30-40kg/m ²
14. Khurana et al, 2017 (218)	Source: Qs Nd:YAG laser Wavelength (nm): 1064, 532 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²): 9.3 (1064nm), 5 (532nm)	<i>In Vivo:</i> Case study, patient with Fusarium solani infection on toe nail	Application of PBM with sequential use of either wavelength cured infection and promoted healthy toe nail growth
15. Wang et al, 2016 (209)	Source: LED array (420), Filtered lamp (540), Diode laser (660, 810). Wavelength (nm): 420, 540, 660, 810 Power (mW): Frequency (Hz): Spot area (cm²): 4	Irradiance (mW/cm²): 16 Time (s): 188 (five times, every 2 days). Energy (J): Radiant exposure (J/cm²): 3	<i>In Vitro:</i> Human adipose-derived stem cells.	Blue and green light were effective in stimulating osteoblast differentiation and increasing intracellular calcium levels than red and near infra-red light. Blue and green light could activate light-gated calcium ion channels.
16. Ashworth et al, 2016 (164)	Source: Wavelength (nm): 450, 510, 660, 860 Power (mW): Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²): Other: 1.93x, 3.85x, 7.70xphotons/cm ² /s	<i>Ex vivo:</i> adapted mouse spinal cord organotypic culture model	Red and near infra-red light are effective antioxidant therapies for spinal cord injury.
17. Merigo et al, 2016 (221)	Source: KTP laser Wavelength (nm): 532 Power (mW): 780 Frequency (Hz): Spot area (cm²): 2.4	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²): 4 Other:	<i>In vitro:</i> Primary bone marrow stromal cells	Green light induces osteogenic differentiation of bone marrow stromal cells.
18. O'Connor et al, 2016 (224)	Source: diode laser Wavelength (nm): 405, 532, 635 Power (mW): 17.5 Frequency (Hz): Spot area (cm²): 1.413	Irradiance (mW/cm²): 12.2 Time (s): 300 Energy (J): 0.0051748 Radiant exposure (J/cm²): 0.003662 Other: Calculations wrong should be 3.66J/cm ² and 5.1748J	<i>In Vivo:</i> C57BL6 mice treated with light and/or Mesenchymal Stem cells	405, 532, and 635 induced increases in mitochondrial activity and reduced apoptosis. Endothelial proliferation increased in response to 635 nm light and combined effects of MSC and the 405 nm wavelength. Reduced TGF-β levels were induced by 532 nm alone and when combined with MSC.
19.	Source: laser	Irradiance (mW/cm²): 50 (630nm and	<i>In Vivo:</i>	All three wavelengths

Fekrazad et al, 2015 (177)	Wavelength (nm): 630, 532, 425 Power (mW): Frequency (Hz): Spot area (cm ²):	532nm), 55 (425nm) Time (s): Energy (J): Radiant exposure (J/cm ²): 2	Diabetes induced male Wistar rats	induced significant increases in wound healing, where red light was most effective.
20. Na, C-S et al, 2015 (223)	Source: laser diodes Wavelength (nm): 532, 658 Power (mW): 30 (532 nm), 60 (658 nm) Frequency (Hz): 20 Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): 180 Energy (J): Radiant exposure (J/cm ²): Other:	<i>In vivo</i> : rat model with induced middle cerebral artery occlusion (MCAO)	Decrease in Bax and cytochrome c levels in hippocampus, increase in hemoglobin, haematocrit, total white blood cell, neutrophil, lymphocyte, monocyte and erythrocyte counts.
21. Burvalev et al, 2014 (168)	Source: Laser (442 nm, 532 nm) and LED (650 nm) Wavelength (nm): 442, 532, 650 Power (mW): 20 Frequency (Hz): Spot area (cm ²): 1.57	Irradiance (mW/cm ²): 30 Time (s): 30-300 Energy (J): Radiant exposure (J/cm ²): 3-31	<i>In Vitro</i> : Mitochondria isolated from rat liver.	Laser of mitochondria at 442nm restored mitochondrial respiration inhibited by NO. Blue light also restored complex IV activity but not complexes I-III. Other wavelengths had no significant effect.
22. Kuboyama et al, 2014 (33)	Source: LED Wavelength (nm): 570 and 940 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): Time (s): Energy (J): Radiant exposure (J/cm ²): 5 (24 sessions) Other:	<i>In Vivo</i> : DBA/1 LacJ male mice with collagen induced arthritis	Reducing swelling induced by both wavelengths. 940nm irradiation induced significant reduction in circulating levels of IL-1β, IL-6 and MMP-3.
23. Cheon et al, 2013 (79)	Source: LED Wavelength (nm): 470, 525, 633 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 3.55 (470 nm), 4.02 (525 nm) and 6.78 (633 nm) Time (s): 3600 (9 days) Energy (J): Radiant exposure (J/cm ²):	<i>In Vivo</i> : Sprague Dawley rats and histological analysis	Blue and green light promoted wound healing significantly. Red light promoted collagen production.
24. De Sousa et al, 2013 (215)	Source: laser (660 nm, and 790nm), LED (700 nm, 530 nm and 460 nm) Wavelength (nm): 660, 790, 700, 530, 460 Power (mW): 60 (660 nm), 50 (790 nm), 15 (700 nm), 8 (530 nm), 22 (460 nm) Frequency (Hz): Spot area (cm ²): 0.03	Irradiance (mW/cm ²): 1911 (660 nm), 1592 (790 nm), 7.46 (700 nm), 8 (530 nm), 22 (460 nm) Time (s): 168 (660 nm), 200 (790 nm), 668 (700 nm), 1250 (530 nm), 456 (460 nm) (every other day, 7 days) Energy (J): Radiant exposure (J/cm ²): 10 Other:	<i>In Vivo</i> : Male Wistar rats wound model and stained for histological evaluation.	530nm, 700nm, 790nm and 660nm induced significant increases in angiogenesis.
25. Tamarova et al, 2009 (205)	Source: LED Wavelength (nm): 480-3400 (range of source, evaluated 'red, orange, yellow, blue, green, violet') Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 40 Time (s): 600 Energy (J): Radiant exposure (J/cm ²): 2.4 (per minute)	<i>In vivo</i> : Male albino rats with area of pain induced by saline injection	Red light was more effective in inducing an analgesic effect. However, all colours induced significant increases in analgesia relative to control.
26. Jackson et al, 2013 (32)	Source: laser diodes (Erchonia GL scanner) Wavelength (nm): 532 nm (6 diodes) Power (mW): 17 per diode, 125 total (sham 1.25) Frequency (Hz): Spot area (cm ²): 516 (target area)	Irradiance (mW/cm ²): Time (s): 900 (two weeks once every 2-3 days). Energy (J): Radiant exposure (J/cm ²): Other:	<i>In Vivo</i> : Human laser irradiation to improve cellulite appearance	532nm improved cellulite appearance on thighs and buttocks.
27. Kim et al, 2013 (219)	Source: Wavelength (nm): 425, 525, 625 Power (mW): Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 6 Time (s): 3600-28800 Energy (J): Radiant exposure (J/cm ²): 21.6-172.8 Other:	<i>In Vitro</i> : <i>Staphylococcus. Aureus</i> , <i>Escherichia. Coli</i> , <i>Porphyromonas. gingivalis</i>	No bactericidal effect induced by red light. Blue and green light were bactericidal where green light also killed <i>S.aureus</i> .
28. Fushimi et al, 2012 (217)	Source: LED Wavelength (nm): 638nm, 456nm, 518nm Power (mW): 2520 (638 nm), 2310 (456 nm), 2500 (518 nm): <i>in vivo</i> . 7560 (638 nm), 6930 (456 nm), 6840 (518 nm): <i>in vitro</i> Frequency (Hz): Spot area (cm ²):	Irradiance (mW/cm ²): 0.25: <i>in vivo</i> 0.75 (638 nm), 0.25 (456 nm), 0.17 (518 nm): <i>in vitro</i> Time (s): 1200 Energy (J): Radiant exposure (J/cm ²): 0.3: <i>in vivo</i> 0.6 (638 nm), 0.3 (456 nm), 0.2 (518 nm). Other:	<i>In Vivo</i> : Mice <i>In Vitro</i> : Fibroblasts and HaCat keratinocytes	Green light decreased wound size. Green and red light accelerated reepithelialisation. Green light induced increases in leptin, IL-8 and VEGF. Keratinocyte migration enhanced by red and green light.
29. Li et al, 2011 (220)	Source: laser (Nd:YAG) Wavelength (nm): 532 nm Power (mW): 40	Irradiance (mW/cm ²): Time (s): 300 Energy (J):	<i>In Vitro</i> : Vascular smooth muscle cells (VMSCs)	Low intensity laser can prevent VMSC proliferation through induction of

	Frequency (Hz): pulsed at 'double frequency' Spot area (cm²): 0.32	Radiant exposure (J/cm²): Other:		increases in markers for apoptosis.
30. De Sousa et al, 2010 (216)	Source: LED Wavelength (nm): 700, 530, 460 Power (mW): 15 (700 nm), 8 (530 nm), 22 (460 nm) Frequency (Hz): Spot area (cm²):	Irradiance (mW/cm²): Time (s): every other day 7 days Energy (J): Radiant exposure (J/cm²): 10 Other:	<i>In Vivo:</i> Wistar rats and fibroblasts grown from biopsy	Green and red light induced significant increases in fibroblast number.
31. Al-Watban et al, 2009 (212)	Source: laser diode Wavelength (nm): 532, 633, 670, 810, 980 Power (mW): 143 (532 nm), 140 (633 nm), 120 (670 nm), 200 (810 nm), 200 (980 nm) Frequency (Hz): Spot area (cm²): 7 (532 nm), 9 (633 nm), 5.25 (670 nm), 9 (810 nm), 9 (980 nm)	Irradiance (mW/cm²): 20.4 (532 nm), 15.56 (633 nm), 22.86 (670 nm), 22.22 (810 nm and 980nm). Time (s): 532nm: 246-1470 633nm: 324-1926 670nm:216-1314 810nm:228-1350 980nm:450-1350 three times per week Energy (J): Radiant exposure (J/cm²): 5, 10, 20, 30 (532 nm, 633 nm, 670 nm, 810) 10, 20, 30 (980 nm) Other:	<i>In Vivo:</i> Wound healing in diabetic Sprague-Dawley rats	PBM accelerated burn healing, particularly visible lasers. Response was dose dependent where highest increase in healing was induced at 30J/cm ² by green light but 20J/cm ² by red light.
32. Tierney and Hanke, 2009 (227)	Source: Diode laser Wavelength (nm): 532 and 940 Power (mW): Frequency (Hz): pulse duration: 60ms (532 nm) and 21ms (940 nm) Spot area (cm²): spot size: 1mm	Irradiance (mW/cm²): Time (s): Energy (J): Radiant exposure (J/cm²): 15 (532 nm) and 100 (940 nm) Other:	<i>In Vivo:</i> RCT, humans with facial telangiectasias	Both wavelengths proved effective in treating facial telangiectasias, but 940nm proved more effective as well as inducing fewer/milder side effects.

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629 4.4 Discussion

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631 It is apparent from this literature review, that whilst the majority of articles reported a positive
632 effect of blue (72%) and green (75%) light PBM, further work is required to demonstrate the
633 importance of the correct recording and reporting of treatment parameters. In fact, in this review
634 69% of all articles failed to report any means of measuring the output of their light source. While a
635 number of studies have highlighted the importance of proper and thorough reporting and recording
636 of treatment parameters (15, 228, 229), it appears these guidelines are yet to be fully implemented
637 in practice. Future efforts are therefore required to ensure the correct reporting of parameters, to
638 enable comparison PBM studies and therefore enable identification of beneficial parameters for
639 therapeutic application.

640 Furthermore this literature review has revealed that for articles reporting the beneficial
641 effects of both, green or blue light application, the majority of publications employed radiant
642 exposures $<10 \text{ J/cm}^2$ (66%, 26/39, excluding articles that did not report radiant exposure values).
643 Interestingly, the beneficial effects of blue and green light included promotion of wound healing
644 (29), reduced inflammation (175), reduction of symptoms in acne (29, 183) and reduced bleeding
645 time following tooth extraction (21) to name a few. The full range of applications of blue and green
646 light found in this review are described in Table 1 and Table 2 respectively. A handful of authors also
647 reported on the biphasic dose response of light (19, 230). For example, *Masson-Meyers et al*
648 investigated the effect of blue light on wound healing *in vitro* using human dermal fibroblast (230).
649 The authors utilised a scratch assay to inflict a 'wound' on cell cultures and following this irradiated
650 cells at 470nm (30 mW/cm^2 , $3\text{-}55 \text{ J/cm}^2$) and evaluated the effect of irradiation on a series of
651 markers for wound healing. The authors reported that at fluence values of 3, 5 and 10 J/cm^2 ,
652 irradiation significantly reduced the secretion of IL-6, a key pro-inflammatory cytokine, increased
653 overall protein production (as a marker for transcription and translational activity) and had no
654 significant impact on wound healing. They also found that irradiation induced mean increases in
655 basic fibroblast growth factor (bFGF) levels, however, this was not significant. Conversely, when
656 utilising a fluency value of 55 J/cm^2 , the authors found that irradiation did significantly reduce rate of
657 wound healing. These data suggest that lower doses of blue light could prove beneficial in inducing
658 decreases in inflammation and promoting gene expression. This theory is in agreement with the
659 'Arndt-Schulz law' in which the application of a stimulus is only beneficial within a relatively narrow
660 therapeutic window. Outside this window a stimulus can either have no effect or induce
661 bioinhibition (231). Interestingly, previous articles have also suggested that 453 nm light is non-toxic
662 up to 500 J/cm^2 , when applied to human skin cells (232). Hence, future work may prioritise the study
663 of the biphasic effect of various wavelengths of blue and green light on cells isolated from different
664 sources in the human body.

665 This review, has also provided evidence for alternative applications of PBM, in which visible
666 light could not only modulate tissue response but also exert antimicrobial properties. Notably the
667 majority of articles citing the antibacterial properties of light use high radiant exposures ($>55 \text{ J/cm}^2$
668 (167, 192, 219)) and these levels could potentially be toxic to eukaryotic cells. Comparatively, de
669 Sousa et al reported that 450 nm light inhibited bacterial growth (*Staphylococcus aureus* and
670 *Pseudomonas aeruginosa*) at doses as low as 6 J/cm^2 (31). Hence, future work may endeavour to
671 determine parameters of visible light required to modulate tissue response whilst also inhibiting
672 bacterial growth.

673 However, it is prudent to highlight one limitation of this review in which radiant exposure
674 values were used to compare literature currently published. Radiant exposure is an important

675 parameter as it takes into consideration a number of other key parameters including irradiance and
676 enables initial establishment of a possible therapeutic window in which blue and green light could
677 induce beneficial effects *in vivo*. However, it is also unreliable as it assumes there is an inverse
678 correlation between both irradiance and exposure time (15). Hence, it is important that authors
679 report all treatment parameters values utilised in studies. This will therefore ensure reliable
680 comparison of current literature and provide further detail as to the parameters that may induce
681 beneficial effects clinically. Future work may also endeavour to evaluate the possible parameter
682 combinations that may induce a beneficial effects within this therapeutic window.

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684 5 Conclusions

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686 This review has provided examples of the wide range of possible targets for various wavelengths of
687 light employed in PBM. These ranged from the application of blue and green light to modulate opsin
688 signalling (17) to the application of red and NIR light to induce cytochrome c oxidase activity (231).
689 We provide evidence for the idea that the majority of these primary mechanisms converge on their
690 ability to modulate ROS production. It has been proposed that small increases in ROS production can
691 induce beneficial effects including increases in cell proliferation, whilst large increases can induce
692 apoptosis signalling pathways (26). Literature currently suggests that light application to 'healthy'
693 cells and tissue induces small increases in ROS production (70), whilst PBM can induce decreases in
694 ROS production in inflamed tissue (111, 233). Hence, PBM could plausibly be applied both as a
695 preventive measure, as well as a means to modulate inflammation in disease. However, further work
696 is required to validate this hypothesis.

697 We also report how PBM induces the activity of downstream signalling pathways, which are
698 modulated by this ROS production. Current literature also demonstrates the wavelength dependent
699 effects of PBM on downstream signalling pathways, where red and NIR light have been proposed to
700 increase the activity of TGF- β signalling (131), whilst blue light has been shown to inhibit the same
701 pathway (234). It will therefore be important in the future to evaluate the wavelength dependent
702 effects of PBM on downstream signalling pathways to provide further indications as to which
703 wavelengths are beneficial for the resolution of different diseases and disorders.

704 This review is also the first report, to our knowledge, which systematically reviews the
705 current literature evaluating the effects of green and blue light PBM both *in vitro* and *in vivo*. We
706 provide evidence that application of blue or green light PBM could have beneficial effects. However,
707 it is apparent that to date, the majority of authors have not appropriately recorded and reported
708 their parameters, meaning that firm conclusions cannot be drawn regarding the optimum
709 parameters to be applied therapeutically.

710 Overall we conclude that PBM exhibits the ability to modulate the activity of an array of
711 signalling pathways, ultimately inducing the beneficial effects seen *in vitro* and *in vivo*. However,
712 further work is required to ensure that experimental studies carry out rigorous spectral
713 characterisation to enable improved reproducibility.

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717 7 *Conflicts of interest*
718 MRH declares the following potential conflicts of interest.
719 Dr Hamblin is on the following Scientific Advisory Boards
720 Transdermal Cap Inc, Cleveland, OH
721 BeWell Global Inc, Wan Chai, Hong Kong
722 Hologenix Inc. Santa Monica, CA
723 LumiThera Inc, Poulsbo, WA
724 Vielight, Toronto, Canada
725 Bright Photomedicine, Sao Paulo, Brazil
726 Quantum Dynamics LLC, Cambridge, MA
727 Global Photon Inc, Bee Cave, TX
728 Medical Coherence, Boston MA
729 NeuroThera, Newark DE
730 JOOVV Inc, Minneapolis-St. Paul MN
731 AIRx Medical, Pleasanton CA
732 FIR Industries, Inc. Ramsey, NJ
733 UVLRx Therapeutics, Oldsmar, FL
734 Ultralux UV Inc, Lansing MI
735 Illumiheal & Petthera, Shoreline, WA
736 MB Lasertherapy, Houston, TX
737 ARRC LED, San Clemente, CA
738 Varuna Biomedical Corp. Incline Village, NV
739 Niraxx Light Therapeutics, Inc, Boston, MA
740 Dr Hamblin has been a consultant for
741 Lexington Int, Boca Raton, FL
742 USHIO Corp, Japan
743 Merck KGaA, Darmstadt, Germany
744 Philips Electronics Nederland B.V.
745 Johnson & Johnson Inc, Philadelphia, PA
746 Sanofi-Aventis Deutschland GmbH, Frankfurt am Main, Germany
747 Dr Hamblin is a stockholder in
748 Global Photon Inc, Bee Cave, TX
749 Mitonix, Newark, DE.

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752 8 *References*

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