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The Use of Immersion Oil in Parasitology Light Microscopic Examination

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ABSTRACT

Aims: to briefly review the history of application immersion oil and specifically in some Parasitology microscopic examination.
Discussion: One way of increasing the optical resolving power of the microscope is to use immersion liquids between the front lens of the objective and the cover slip. The application of immersion oil in microscopic examination began in the mid-17th century. The principle of immersion oil in microscopic examination is to improve the resolution and clarity of the image by reducing the refractive index mismatch between the specimen and the objective lens. In the context of micro-sized parasitic agent, adding this oil will greatly enhanced visualization and in turn will have a direct impact on improving the characteristic morphological image/appearance, thereby enabling observers to optimize microscopic assessment for parasitological examination
Conclusion: The addition of immersion oil in Paarasitology microscopic examination is surely improve optical resolution and clarity of the image and positively support Parasitologist to make correct diagnosis

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Keywords: resolution, magnification, lens, numerical aperture object, micro-parasite, morphology

1. INTRODUCTION

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Light microscopy is a powerful tool for examining micrometric objects (unseen by naked eyes) [1,2], organic (motile or alive) or non-organic (non-motile or in-organic) [3], across a wide range of applications in systems Biology [4] by providing an enlarged image of small structure that deliberately interact with visible light [5], for example their light-absorption, light-reflection and even light-scattering. Light microscopy is used to make small structures and samples visible by using the aid of light, naturally or electrically [1-5]. Contemporary light microscopes are able to magnify objects up to about a thousand times. Since most cells are between 1 and 100 μm in diameter, they can be observed by light microscopy, as can some of the larger subcellular organelles, such as the nucleus, cell membrane, cytoplasm, chloroplasts, mitochondria and cell wall surroundings [6].
This simple microscopic approach is useful to help observer to (1) get the larger picture of the object for identification, (2) analyze and understand the morphologic appearance of the sample for differentiation and even perhaps (3) to explore the material construction in more

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35 detail. Light microscope also allows examiners to observe sophisticated scrutiny of the
36 microscopic world, such as how substances diffuse across a cell membrane [1].
37 Immersion oil is used mainly in high magnification of light microscopic examination to improve
38 the resolution and clarity of the object being observed [7]. Parasitology is very blessed with
39 the existence of microscopic examinations, especially those that use immersion oil, because
40 it helps examiners identify microscopic objects correctly. The aim of this mini review is to reveal
41 the function of immersion oil in Parasitology examination using light microscope and how it
42 works.

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44 2. HISTORY OF IMMERSION OIL

45 The application immersion oil cannot be separated by the invention of microscope. It is
46 believed that two Dutch spectacle-collaborators and also father-and-son team, Hans and
47 Zacharias Janssen, create the first microscope in 1590; they discovered that, if they
48 intentionally put a lens at the top and bottom of a tube and looked through it, objects observed
49 on the other end of the lens surprisingly became aggrandized [8]. Then in 1667 or 1668, Robert
50 Hooke's well-known presentation, "Lectures and Collections" which was released scientifically
51 in the same year with his masterpiece "Micrographia", which delineates Hooke's extensive
52 analysis using his simple microscope. Hooke was actually the first to suggest the technique of
53 Immersion. He addresses: "that if you would have a microscope with one single refraction,
54 and consequently capable of the greatest clearness and brightness, spread a little of the fluid
55 to be examined on a glass plate, bring this under one of the globules, and then move it gently
56 upward till the fluid touches and adheres to the globule".

57 By the year 1812, a breakthrough idea of Brewster suggested the application of immersion of
58 the objective into the liquid medium [11] while in almost that same time, Giovanni Battista
59 Amici began pointing the difficult situation regarding chromatic aberration [12] and in 1840, he
60 succeeded making the first working immersion lenses, which initially were designed to be used
61 with oils having the same refraction as glass, homogeneous-immersion [13]. This equipment
62 was far from perfect.

63 Robert B. Tolles, a genius considered as one among three American Microscope pioneer and
64 builder [14], has a brilliant idea; his insight was increasing the apertures that improves
65 revelation [15]. Amici's main focus was aimed to achieve re-correction from a common
66 condition found in optics named aberrations, a property of optical systems (e.g., lenses), that
67 causes light to be spread out over some region of space rather than focused to a point [16, 17].

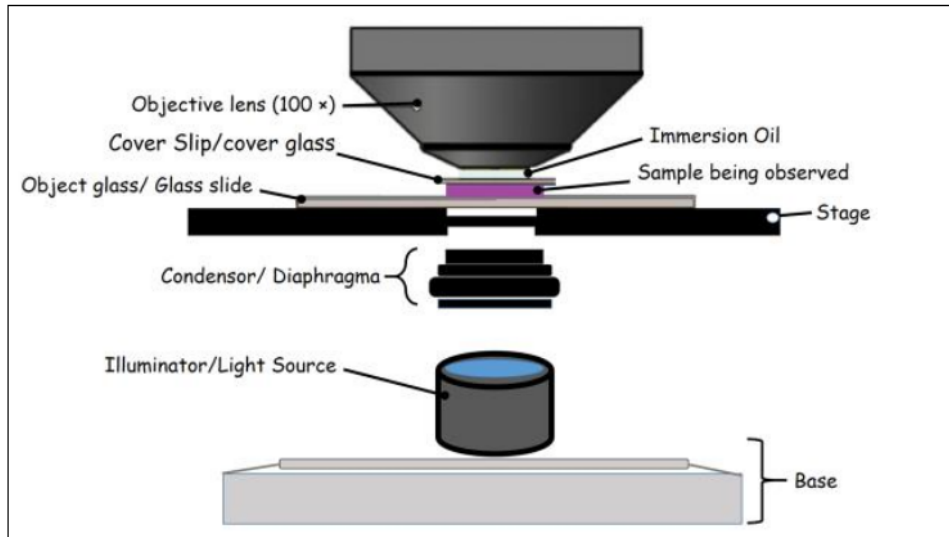
68 The principal limitation of the light microscope is its resolving power [5,17-20]. Resolving
69 power of a microscope is a function of refractive index [21-23]. The resolution limit of a
70 microscope is the shortest distance between two nearby objects when the images formed by
71 the microscope are properly differentiated. The resolving power of the microscope is $X_{min} =$
72 $1.22\lambda / \text{numerical aperture}$. Unfortunately, frontiersman microscopists were hindered by three
73 common obstacles, namely (1) optical aberration that unfortunately occur because of lens
74 curvature and dispersion [17], (2) unclear and blurred images due to incorrect use [18], and
75 (3) poor lens design- commonly due to low quality and of imperfect shape, producing sub-
76 optimal imaging [19], which impeded high-resolution inspections until late in the industrialized
77 era [20]. Aberrations were partially able to be corrected by the mid-19th Century, thanks to the
78 introduction of achromatic objectives that reduced chromatic aberration [19] and improvised
79 of numerical apertures [21] to around range from 0.04 for a 1 X objective to 0.95 for a 60 X
80 objective (the largest possible NA) for dry objectives [22] and up to 1.25 for homogeneous
81 immersion objectives for most research grade microscopes condenser lens systems (1000x
82 magnification) [23]. The next section will discuss the principles and physical properties of

83 immersion oil that commonly use in light microscope that will help augment the size of object
84 being analyzed.

85 3. IMMERSION OIL: ITS PRINCIPLES AND PHYSICAL PROPERTIES

86 ² In light microscopy, oil immersion is a technique used to increase the resolving power of a
87 modern microscope [7]. The application of this oil are aimed to (1) increase the resolution
88 (finer resolution) and (2) improve clarity (brightness) of microscopic image [24]. These two
89 distinctive characteristics are most crucial below high magnification; so it is only the higher
90 power, short focus, objectives that are usually designed for oil immersion [25].

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93 Fig. 1. Schematic of anterior aspect of regular light microscope with the application of
94 immersion oil

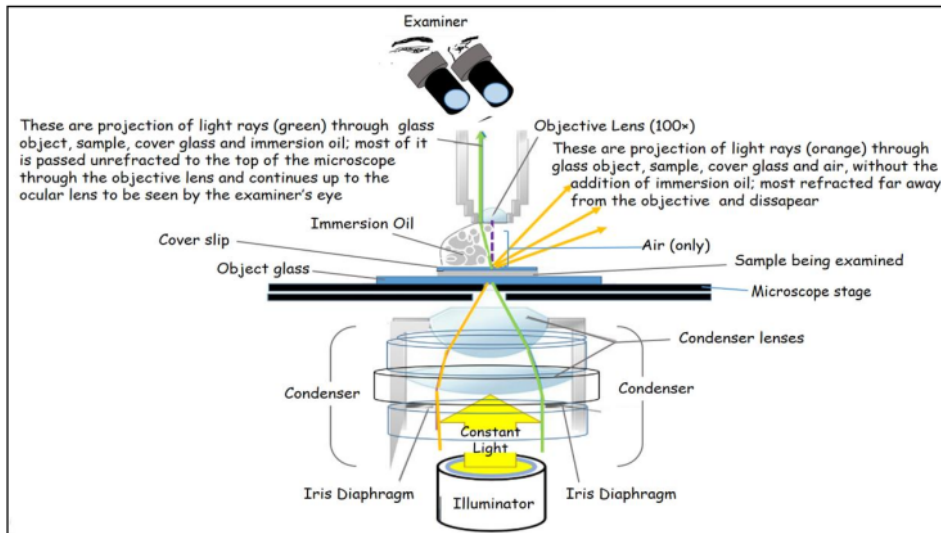
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96 ² The application of this oil is between the front lens of the objective and the cover slip
97 [11,24,25]. Most objectives in the magnification range between 60x and 100x (and higher) are
98 designed for use with immersion oil [26]; and by doing so thereby increasing the numerical
99 aperture of the objective lens. Technically, a gain in resolution by a factor of about 1.518 is
100 attained when immersion oil is substituted for air as the imaging medium. The factor (1.518)
101 which is very close to the refractive index of glass. All reflections on the path from the object
102 to the objective are eliminated in this way.

103 The oil used must be transparent [24] with high refractive index and It works based on the
104 principle of matching the refractive index between the specimen and the microscope objective
105 lens. By placing a drop of immersion oil between the lens and the specimen, the light rays
106 passing through the specimen are less likely to scatter, resulting in a sharper image [26]. The
107 immersion oil also helps to reduce the loss of light due to refraction at the air-glass interface
108 [25]. This allows for higher magnification and better visualization of fine details in the specimen
109 [11]. The properties of the immersion oil, such as refractive index, viscosity, and erodibility,
110 are important factors in its effectiveness for microscopy [21-23].

111 ¹ The important key of objective lens relies on the beam of light emitted from illuminator that
112 pass through series of media from air into glass (condenser lens) and then into object glass,

113 specimen being analyzed and then cover slip with its mounting and then immersion oil or air
 114 (if no immersion oil used). The direction of the beam is bent and when it passes back from
 115 glass to air it is bent back again to its initial, original direction, and this surely impinge with high
 116 power lenses because of limited the amount of light which actually can reach and enter the
 117 lens, affect the numerical aperture (NA) of the lens and consequently its resolving power. To
 118 overcome this bending effect, adding oil to replace the air between the specimen and the lens
 119 which has the same refraction index as glass, will surely make the light passes in a straight
 120 line from glass through the oil and back to glass as though it were passing through glass all
 121 way (see fig. 1 and fig. 2) [27]



122 Fig. 2. Schematic representation of the difference of route of refraction, when light pass
 123 through immersion oil (left side) and air (right side); objective lens use is 100x magnification.
 124 Immersion oil is used in an optical/compound microscope during microscopy analysis in order
 125 to boost the resolution of the microscope. It is commonly applied in simultaneity with a high-
 126 resolution objective lens (100x magnification) to increase the numerical aperture of the lens,
 127 which allows it to gather more light and produce a clearer, higher-resolution image. Immersion
 128 oil is placed between the objective lens and the specimen, and it has a refractive index that is
 129 very close to that of the glass used in the objective lens. This allows the light to pass through
 130 the oil and the glass without being refracted too much, which helps to improve the resolution
 131 of the microscope.
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133

134 Adding Oil immersion in dry lenses actually useless and always changed/deformed the actual
 135 images or even distorts it [17]. That is why a complete cleansing of the slide and also the
 136 objective lens after using an oil immersion from previous analysis is a must, before using a
 137 dry or air objective lens (*i.e.* lens with magnification 5x, 10x or 40x) to check or image it again.
 138 The use of xylene to remove (clean) immersion oil from microscope slides after examination
 139 for acid-fast bacilli have been reported [28]. By Cleaning the entire left over immersion oil from
 140 previous application also circumvents tainting the air objective with an improper substance
 141 which is not needed [29,30]. So for every microscopists, maintaining the cleanliness of
 142 microscope optics is pivotal to ensure the readiness to use and guarantee high-quality imaging
 143 [31]. Micro particle such as dust, fingerprints, excess immersion oil, or mounting medium on

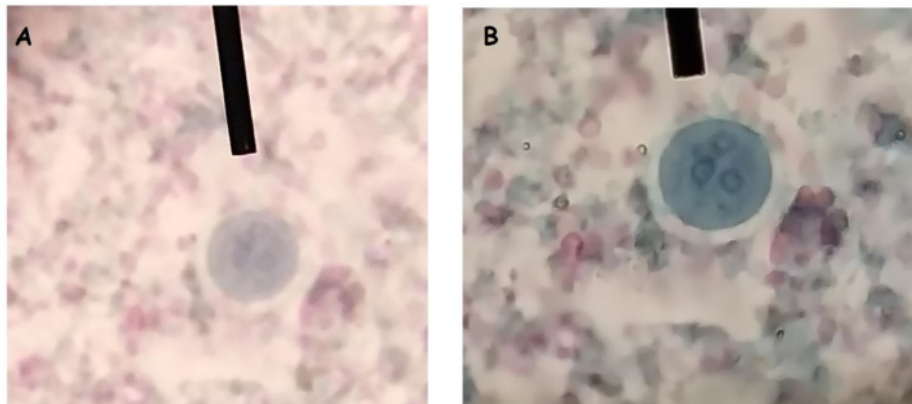
144 or in a microscope causes reduction in contrast and resolution and all of these will directly
145 affecting its imaging performance; especially the lens surfaces which is very fragile and sensitive
146 to scratches, stains and contamination [29-31].

147 4. APPLICATION OF IMMERSION OIL IN PARASITOLOGY MICROSCOPIC 148 ANALYSIS

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150 If something suspicious is seen by the examiner, a higher magnification may be necessary,
151 and this is the ultimate purpose of using immersion oil in an optical microscope during a
152 stained or wet-mount Parasitology microscopic analysis.

153 Most parasitic protozoan in humans are small and within range $<50\ \mu\text{m}$ in size [32]. The
154 smallest (predominantly as an intracellular forms) are $1\text{-}10\ \mu\text{m}$ in diameter or in its longest,
155 but with the exception of *Balantidium coli* may reach up to $150\ \mu\text{m}$ [33]. Protozoa are a
156 polyphyletic group of mono-celled eukaryotes that are actually heterotrophic, self-motile, and
157 lack cell walls [32]. Due to its smaller size, compared to helminths or other type of ectoparasite,
158 protozoan parasites, especially in the intracellular form, present a bigger dispute to make a
159 correct diagnosis based on microscopic examination [34,35].

160 Globally known small-size disease-causing protozoan parasites include *Plasmodium* spp
161 ranging from $1\text{ to }20\ \mu\text{m}$ in size depend on their stage [36], *Entamoeba histolytica* usually
162 measure range $10\text{ to }15\ \mu\text{m}$ (cyst form) and $20\text{ to }30\ \mu\text{m}$ (cyst form) [37], *Giardia lamblia*
163 whose trophozoites resemble a pear or teardrop and measure $10\text{-}20\ \mu\text{m}$ in length, $5\text{-}15\ \mu\text{m}$ in
164 width and $1\text{-}2\ \mu\text{m}$ thickness [38], *Microsporidia* spp whose resistant spore usually measure
165 $1\text{--}4\ \mu\text{m}$ [39], *Cryptosporidium parvum* with its oocysts are measure $4\text{ to }6\ \mu\text{m}$ in diameter
166 [40], *Trichomonas vaginalis* with averages $26\ (21\text{-}32)\ \mu\text{m}$ in total length, with $9.5\ (7.4\text{-}11.4)$
167 μm of body length and $6.8\ (5.3\text{-}7.7)\ \mu\text{m}$ of width [41], *Toxoplasma* spp whose bradyzoites are
168 $7 \times 1.5\ \mu\text{m}$ [42], *Trypanosoma* spp whose length is about $25\ \mu\text{m}$ in length for *T. brucei* [43],
169 *Babesia* spp with an average size of $4.5 \times 2\ \mu\text{m}$ [44], *Leishmania* spp whose amastigotes
170 appear round in shape and $2\text{--}4\ \mu\text{m}$ in diameter while cultured promastigotes range between
171 $15\text{ and }25\ \mu\text{m}$ in length and are ellipsoid to slender in shape [45], *Cyclospora cayatanensis*
172 which oocyst very consistent in shape (round or slightly flattened on one side) and size range
173 only between $8\text{-}10\ \mu\text{m}$ [46] and *Blastocystis hominis* which size varies from $6\text{ to }40\ \mu\text{m}$ [47].



174
175 Fig 3. *Entamoeba histolytica*, Cyst form with four nuclei (total magnification $1000\times$ in Olympus
176 CX21LED). A. without Immersion oil, B. with Immersion oil. Photo taken manually using phone
177 cell with both using 3.6 optical zoom. There is a clear difference on clarity and brightness of
178 the image
179

180 Those long lists of previously mentioned small size parasites revealed the complexity of
181 microscopy examination aimed to locate and then to pin point its exact position and followed
182 by magnify the parasite, so that the examiner can confidently determine the typical
183 morphological characteristics, e.g., the species, stage/stadium as well as its density.
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186 **5. CONCLUSION**

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188 In daily and routine light microscopy use in the laboratory, the goal of adding oil immersion on
189 the sample's cover glass is simply to improve the resolving power of a microscope. This goal
190 is accomplished by immersing both the objective lens and the specimen in a transparent oil of
191 high refractive index, thereby increasing the numerical aperture of the objective lens; all of
192 these means better visualization in size and clarity. In Parasitology, several parasites are
193 actually very small (e.g., *Plasmodium* spp) and in order to visualize them properly, the slide
194 must be analyzing in higher magnification (objective lens 100x) with the addition of a drop of
195 immersion oil.
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198

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204 **COMPETING INTERESTS**

205

206 "Authors have declared that no competing interests exist."
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208 **AUTHORS' CONTRIBUTIONS**

209

210 " 'Author FES' designed the study, performed the analysis, conducted the phone cell
211 photography, wrote the protocol, and wrote the first draft until final manuscript."
212

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213 **CONSENT (WHERE EVER APPLICABLE)**

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215 Not needed
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218 **ETHICAL APPROVAL (WHERE EVER APPLICABLE)**

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220 Not needed
221

221

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