Zoological and Biological Role of Parasites, Pathogens, Worms in Food Web and Future Perspectives

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Abstract

Instructive approaches for testing emerging malaria vaccine candidates could potentially have a lot of translational value. The potential of genetically altered Plasmodium species that do not grow beyond the liver stages, in triggering a protective immunity. Giardia-lambila, Ascaris-lumbricoideis, Trichuris-trichiura, Schistosoma species and Entamoeba-histolytica are most intestinal worm parasites around worldwide. Amebiasis is caused by Entamoeba histolytica invading the intestinal mucosa and causing severe mortality and morbidity. Babesiosis is tick borne infectious disease which causes anaemia, fever, haemoglobinuria, and frailty in the animals. Various bacterial species have been found in fish’s natural habitats. Bacterial colonies can be seen on fish’s gills and skin as a result of persistent contact to unclean water, whereas polluted water or food might harm the gastrointestinal tract. Salmonella nonstphoidalis is the most common zoonotic bacterial food borne infection in humans. Salmonellae are common around the world, and they are the most common pathogenic bacteria in both humans and animals. Engineering the genes which code for pathogenicity effectors and host adaptability could be the key to completely identifying the causal relationship among a gene deficiency and infections in organoids.

Keywords: Immunity, zoonotic bacterial, food borne infection, pathogenicity effectors.

INTRODUCTION

Instructive approaches for testing emerging malaria vaccine candidates could potentially have a lot of translational value. A vaccine to completely prevent the malaria need for its eradication. Malaria vaccines are the subject of two additions to this RT. Examples, strategies, and present situation of carbohydrate antigen-based vaccines targeting malaria, leishmaniasis and toxoplasmosis . In malaria studies, the concept of a live attenuated vaccine has a lengthy history. Parasites that have been treated or pharmacological attenuated have been employed; however they may not be the best option. The potential of genetically altered Plasmodium species that do not grow beyond the liver stages, in triggering a protective immunity. They believe that achieving the parasite’s potential will necessitate substantial investment in the murine models in order to improve its translational relevancy [1-7]. Intestinal-parasitic-infections (IPI) are generated by intestinal-helmiths and protozoans, and they continue to be one of the most potentially serious issues in underdeveloped nations with inadequate water supply and sanitation [1, 2]. Approximately 3.5 billion of people worldwide infected, with 450 million becoming unwell as a consequence of one or more gastrointestinal parasite infections [3]. Hookworms, Giardia-lambila, Ascaris-lumbricoideis, Trichuris-trichiura, Schistosoma species and Entamoeba-histolytica are most intestinal worm parasites, with more than 10.5 million new cases recorded yearly [4]. Malnutrition, anaemia, diarrhoea, intestinal blockage,
physically and mentally stunted growth, malabsorption, poor work ability, and a slower growth rate are all possible consequences of these diseases [5, 6]. Intestinal parasitic infections are very common amongst some of poor and are linked to lower house income, poor individual and situational hygiene, overpopulation, restricted access to clean drinking water, a tropical climate, and lower altitude [7-9].

Only a few nations in Sub-Saharan Africans have documented parasite species infecting the economically important marine fishes, thanks to coordinated efforts by indigenous parasitologists (e.g. Senegal, South Africa, Nigeria). The majority of these are taxonomy records or broad examinations of parasitic faunas connected with marine hosts, that might or might not be commercially valuable. In most sections of Sub-Saharan Africans, limited to no multidisciplinary investigation is undertaken, thus parasitological findings are rarely used to assist fishery management practices. The present state of parasitological investigations on economically valuable marine fishes in Sub-Saharan Africans is summarized in this article [10-12].

Amebiasis is caused by the cytolytic protozoan parasite (Entamoeba histolytica) invading the intestinal mucosa and causing severe mortality and morbidity on a worldwide scale. A review was conducted to identify the processes whereby the parasite induces aggressive disease is provided, such as the historical background, epidemiological studies, and pathophysiology of amebiasis, parasitic organism biology, in vitro and in vivo studies of pathogenicity, host’s immune reactions, and preventative health and biologic control options. Novel findings into E. histolytica molecular genetics, particularly its adhesion and cytolytic capabilities, have opened the door to various new techniques to bio control of this invasive, devastating parasitic disease [11-14].

**Zoological and biological role of parasites, pathogens, worms in food web**

Gut dysbiosis, or an imbalance in composition of intestinal-microbiome, is one cause of inflammaging. Decline of immunostimulatory microbes species can result from this imbalance, that is aggravated by pro-dysbiotic components of contemporary living, like antibiotic use and the so called Western-diet. The old friends’ idea is one view of a cause of these dysbiotic consequences. This theory claims that the person’s immune system developed to function optimally in a dirty world with more bacteria and helminth parasite, whose elimination causes pathological immunological hyperactivity. Prior to age, gut dysbiosis does have a role in allergy and auto-immune inflammation. Recovery of old friend species should decrease such inflammation hyperfunction and lessen disease, according to one conclusion of old friends argument. Likewise, fecal microbiota transplantation from healthy individuals has been proposed as a possible anti-inflammatory treatment. Whereas the microbiome’s potential function in inflammaging has piqued biogerontologists’ interest, little attention has been paid to macrobiome's potential role in ageing, particularly helminth parasite such as tapeworms, nematodes and flukes [12, 13].

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**Fig-1: Shows role of Zoological and biological manifestations of parasites, pathogens, worms**

**Babesiosis and other infections**

Babesiosis is tick borne infectious disease which causes anaemia, fever, haemoglobinuria, and frailty in the animals. Piroplasmosis, red water, Texas fever, tick fever, tristeza and bovine babesiosis are some of the other names for the disease [5]. Theileriosis in goats, cattle, sheeps, and in captive and wild ungulates is caused by the Theileria spp. [10].
Theileriosis is a haemoparasitic illness caused by the Apicomplexa (a Theileriosis protozoa). Domesticated and wild ruminants are affected by Theileria spp, which are found primarily in Europe, Asia, Australia, and Africa [17-19]. Ticks are the carriers of parasites. These parasites go through a cycle of merogony within lymphocytes, eventually producing micro-sopic merozoites that penetrate red cells and turn into plasmoids. Theileriosis is spread by a number of t- vectors, that can induce infections that vary from clinically insignificant to deadly [9]. The infection is also a haemoparasitic illness caused by the Babesia- protozoa (Phylum: Apicomplexa), that primarily affects ruminants [6]. The injection of sporozoite parasite in to blood-stream during the ingestion of a blood meal starts the disease of vertebrate host [14-16].

Cryptosporidiosis is a zoonotic parasitic infectious disease caused by the intestinal parasite Cryptosporidium-parvum and spreads through food and water contamination. Infection is contracted by consuming food or water infected by Cryptosporidium species oocysts, that can be lethal in some situations [27]. In 2007, a research was carried in Zoo Negara (national Malaysian) to re-evaluate the presence of Cryptosporidium parasites in samples taken from birds . Reported in this study, they discovered a link between cryptosporidiosis and the infected birds' handlers, confirming zoonotic propagation. In 2007, a cross sectional research was carried out between look into the distribution of Giardia and Cryptosporidium species. Cryptosporidiosis was found to be prevalent in 4.1 percent of Orang Asli in Malaysia, without any gender bias among Giardia and Cryptosporidium species, demonstrating that such infections remain serious health issues among Orang Asli. In 2012, a research was carried in two urban hospitals in Selangor state to evaluate the incidence of cryptosporidiosis in 130 diarrhoeal diseases samples of children. It showed that 4.62 percent of samples tested positive for the Cryptosporidium species., showing that hygienic standards in childcare centers, where several parents abandon their kids for long periods of time, must be maintained and checked. A cross sectional research of 388 migrant-workers in Peninsular Malaysia was done in 2017 to investigate the prevalence of protozoan infections. Microscopy testing revealed 42 (10.8 percent) Giardia species positive samples and 12 (3.1 percent) Cryptosporidium species positive samples, highlighting the potential dangers of water-borne and food-borne infections in the spreading of such diseases. Cryptosporidiosis has the ability to spread throughout and become a major health issue. It can be fatal, especially among older, young kids, and those with AIDS. Proper monitoring of food and water supplies utilizing investigative approaches like Microbial Source Tracking (MST) can thereby differentiate between animal and human source of fecal pollution, limiting its onset [15-20].

Various bacterial species have been found in fish’s natural habitats. Bacterial colonies can be seen on fish’s gills and skin as a result of persistent contact to unclean water, whereas polluted water or food might harm the gastrointestinal tract. When immunologic resistance is weakened, contamination of fish’s muscles is also a concern. On average, only a few microbes can be detected on fish skin. Total bacterium count (TBC) on the salmon skin (Salmo salar) was found to range from 102-103 CFU/cm2. Conversely, a research in Turkey found a larger number of 101 to 107 CFU/cm2 on the skin of salmon, with aerobic bacteria found more frequently than anaerobic microbes. It is commonly acknowledged that bacteria found on the skin of fish are similar as those present in the polluted water, including these genera and species as Aeromonas species (Aeromonas bestiurum, A. schubertii, A. caviae, A. jandaei, A. hydrophila, A. veronii), Flexibacter species, Psychrobacter species, Proteus species, Providencia species, Pseudomonas fluorescens, Moraxella species, Acinetobacter johnsonii, Enterobacter aerogenes, Vibrio fluvialis, Micrococcus luteus, Alcaligenes piechaudii, and Escherichia coli [21, 22].

In the literature, there is substantial debate about how the parasite degree differs from those of free living animals. Because parasites are often highly specialized parasites may have a lower in degree (number of prey per hosts) than predators that live in the wild. Nevertheless, depending on how the measurement is done, parasites have been found to increase total connectance. This implies that parasites have more connections on average than their free living counterparts. Concurrent predation will increase the I - degree of free living predators and the out degree (number of predators) of parasites, affecting degree. These findings point to degree as a physically differentiating trait of parasitic species, especially when in degree and out degree are taken into account individually [23, 24].

Subgroups and stability are common concepts in both domains, so it seems obvious that they should readily come together. However, despite active intervention on the both sides [2, 4], this may not have accomplished. Because the two fields are heading in opposite directions, the scenario is unlikely to improve. Furthermore, the food web approach has a basic weakness that drastically hinders the parasitological approach. The fundamental issue is that any analyses' results only as effective as its source, and also most food web analysis' raw data are imprecise and often erroneous descriptions of nature [25, 26].

Microscopy is commonly employed, although it only identifies eggs of the Taeniidae family (which comprises Echinococcus species and Taenia species)
without genus or species separation. In one review, 12 research were documented in which eggs of Taeniidae were detected on fresh produces. Various techniques have been used, but most were focused primarily on elution from fresh produce in salt solution, often with emulsifiers like 0.1 percent Tween-80 or 1 percent sodium dodecyl sulphate, accompanied by concentration, by either sedimentation (some with centrifugation) or sucrose flotation or formal ether concentration, accompanied by microscopic examination. Critical information such as the detection limit and reproducibility were missing from all of these papers. There had been no attempt to investigate validation or to add internally or externally quality control samples or data [27, 28].

The majority of sediment and soil food webs are heterotrophic (i.e., rely on the allochtonous or external input of organic compounds like those which are taken from the animal or plant matter). Despite the fact that shallower intertidal food-webs have strong primary production, organic material mineralization typically outpaces local producers because organic material from the surrounding water serves an additional organic source for benthic food-web. Chemoautotrophy is used to maintain some extremely confined deep sea habitats, like hydrothermal vents and cold seeps, by oxidizing reduced chemicals from the sea bottom. The great bulk of the deep ocean, however, is capable of photosynthesis [29-30].

Salmonella nontyphoidalis is the most common zoonotic bacterial food borne infection in humans. Salmonellae are common around the world, and they are the most common pathogenic bacteria in both humans and animals. They are the most commonly bacterial isolates of food borne infectious diseases, accounting for approximately 93.8 million food borne diseases and 155,000 fatalities per year globally [4]. Salmonella has been identified as a leading source of food borne illness and a serious public health issue around the world, with rising concerns about the introduction and transmission of antimicrobial resistant forms [12], particularly in developed nations. Salmonella infections in animals and humans that are resistant to antibiotics are a worldwide issue, especially in developing nations [28-33].

**CONCLUSION**

Microorganisms have a variety of interconnected virulence features that are continually moving toward the formation of infections, triggering disease, and allowing them to remain inside the host. Pathogen-effector research could lead to the creation of new quick diagnosis tools and detection methods, treatment medicines, and vaccinations to help control the food-borne infections. Organoids are establishing the door for more fruitful studies of FBP-virulence at molecular scale. Engineering the genes which code for pathogenicity effectors and host adaptability could be the key to completely identifying the causal relationship among a gene deficiency and infections in organisms. The *S. enterica Typhimurium* invasion cycle utilizing enteroids, and discovered innovations concerning the involvement of recognized virulence determinants.

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