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DEVELOPMENT OF A PORTABLE INCUBATOR FOR THE
DETECTION OF COLIFORM IN WATER USING IOT

BY

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¹
A PROJECT REPORT SUBMITTED TO THE DEPARTMENT OF
ELECTRICAL & INFORMATION ENGINEERING, IN PARTIAL
FULFILLMENT OF THE REQUIREMENTS FOR THE AWARD OF
THE BACHELOR OF ENGINEERING DEGREE IN ELECTRICAL
AND ELECTRONICS ENGINEERING.

SUPERVISOR:

DR. OSEMWEGLI OMORUYI

JUNE 2023

DECLARATION

I, OLONISAKIN DAVID AKOLADE (18CK024244), hereby declare that I carried out the work reported in this project in the Department of Electrical and Information Engineering, Covenant University, under the supervision of Dr. Osemwiegie Omoruyi. I also solemnly declare that to the best of my knowledge; no part of this report has been submitted here or elsewhere in a previous application for the award of a degree. All sources of knowledge used have been duly acknowledged.

.....
OLONISAKIN DAVID AKOLADE

18CK024244

CERTIFICATION

This is to certify that the project titled "**DEVELOPMENT OF A PORTABLE INCUBATOR FOR THE DETECTION OF COLIFORM IN WATER USING IOT**" by OLONISAKIN DAVID AKOLADE, meets the requirements and regulations governing the award of the Bachelor of Engineering in Electrical and Electronics Engineering degree of Covenant University and is approved for its contribution to knowledge and literary presentation.

Dr. Osemwiegie Omoruyi

(Supervisor) Sign: _____ Date: _____

Internal Examiner Sign: _____ Date: _____

Dr. Isaac Samuel Sign: _____ Date: _____

(Head of Department)

DEDICATION

This research project is dedicated to my parents, Dr. Rotimi Olonisakin and Mrs. Bolatito Olonisakin who were very understanding and supportive throughout the time taken to complete this project, providing the resources necessary to make this project possible. I also dedicate this project to the countless communities in Nigeria who lack access to clean water.

ACKNOWLEDGEMENTS

10 First and foremost, I would like to begin by expressing my deepest gratitude to God for His divine guidance, wisdom, and grace throughout this project. I am forever grateful for His constant presence and blessings in my life.

5 I would also like to extend my sincere appreciation to my parents for their unwavering love, support, and encouragement. Their sacrifices, guidance, and prayers have been a source of strength and motivation throughout this journey. I am truly blessed to have them as my role models.

37 I would like to take this opportunity to express my deepest appreciation and gratitude to my exceptional project partner, Oguibe Favour Ozioma. Together, with a harmonious blend of teamwork, unwavering dedication, and the power of prayers, we successfully transformed our shared idea into a remarkable reality.

39 I would like to express my heartfelt thanks to my project supervisor, Dr. Osemwegie Omoruyi for His invaluable guidance, expertise, and support. His insightful feedback, constructive criticism, and dedication have been instrumental in shaping the direction and success of this project.

Furthermore, I want to thank the Head of Department, Prof. Isaac Samuel, and all my lecturers for impacting me with knowledge and passing me through various tasks that have, in one way or the other, prepared me for the gallant execution of this project.

2 Finally, I would also like to acknowledge the support and encouragement of my lovely sisters, Favour and Delight Olonisakin, who have been there for me through thick and thin. Their kind words, motivational messages, and belief in my abilities have helped me to stay focused and motivated.

ABSTRACT

75

Having access to clean and safe drinking water is considered a crucial human right, yet millions of individuals continue to live without it. The consumption of water that has been contaminated can increase the risk of experiencing outbreaks of different diseases that are transmitted through water, which are a leading cause of death worldwide. This project aims to develop a portable and low-cost incubator for detecting coliform bacteria in water, which is achieved by using a light-dependent resistor (LDR) and a heating element controlled by an IOT equipped microcontroller (ESP32). The incubator creates a controlled environment for water samples to incubate and for the LDR sensor to detect changes in turbidity, indicating the presence of coliform bacteria. Additionally, a Digital Humidity and Temperature (DHT22) sensor and a heating element helps maintain a temperature range of 35-37°C, the ideal temperature for optimal coliform bacteria growth. The incubator also features a Liquid Crystal Display (LCD) displaying resistance and temperature readings. After the data is collected, it is transferred to the microcontroller and subsequently transmitted to the Blynk application for continuous monitoring and analysis in real-time. The portability and affordability of this incubator makes it appropriate for use in remote areas where access to laboratory equipment is limited. This study carries significant implications for the advancement of low-cost and portable water quality monitoring systems, potentially leading to improvements in public health.

Keywords: *portable incubator, coliform bacteria , water monitoring, light-dependent resistor.*

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CHAPTER ONE

INTRODUCTION

1.1 Background of Study

The research was inspired by the need for precise and trustworthy techniques for identifying waterborne infections, especially coliform bacteria, in places with few laboratory facilities. Coliform bacteria are used as a quality gauge for water and are a sign that potentially dangerous diseases may be present. Ensuring the availability of safe drinking water stands as a key priority in public health efforts because it is intimately related to human health. Consuming water that contains hazardous chemicals or pathogenic organisms has a major influence on one's health[1].

The lack of low-cost, simple-to-use, and accurate techniques for remote water monitoring and coliform detection is the key issue driving this research. Due to difficulties with transportation, sample storage, and inadequate laboratory facilities, traditional laboratory-based approaches are frequently not practical in remote places. Thus, the creation of a mobile incubator for coliform detection could be the answer to this issue.

The project is aimed towards developing a portable incubator that can quickly and accurately detect coliform bacteria in water using an approach that is both affordable and simple to use. One approach to detect these organisms involves filtering a 100 mL water sample through a membrane. Subsequently, the filtered membrane is incubated on specialized media at temperatures ranging from 35 to 37 °C for E. coli or 44 to 45 °C for TTC[2]. The issue of restricted access to laboratory facilities for water monitoring and coliform detection can be resolved by the portable incubator, especially

in distant locations. The study is important because it may increase access to clean water, especially in locations where waterborne diseases are common. The mobile incubator might also be used for environmental research and monitoring, which would improve water management and public health.

1.2 Significance of Study

The development of a portable incubator for remote water monitoring and detection of coliform is an important initiative that can improve access to safe water, particularly in remote areas. Traditional laboratory-based methods for coliform detection are often not feasible in such areas due to transportation, sample storage, and limited laboratory facilities.

This project aims to address the problem of limited access to laboratory facilities by providing a low-cost, easy-to-use, and reliable method for remote water monitoring and coliform detection. The portable incubator is designed to use a LDR and a light bulb, where a resistance drop or increase is detected depending on the coliform growth. Afterward, this data is transmitted to the ESP32 for additional analysis. Additionally, a DHT sensor is in place to regulate the temperature of the incubator at 35-37 degrees Celsius via the heater. The significance of this study is highlighted below;

1. Improve access to safe water, particularly in remote areas.
2. Enhance public health efforts in identifying and preventing waterborne diseases.
3. Contribute to water management and environmental monitoring research.
4. Increase efficiency in water monitoring by eliminating transportation of water samples for analysis in a laboratory

5. Enable citizen science initiatives for community-led efforts towards sustainable water management.

1.3 Problem Statement

The current methods for detecting coliform bacteria in water samples require laboratory analysis, which can be time-consuming and resource-intensive. This can limit the ability to effectively monitor water quality, particularly in remote areas. Therefore, there is a need for a portable incubator that can detect coliform bacteria remotely, and enable more efficient and timely water monitoring activities.

1.4 Aim and Objectives

Aim: To develop a portable incubator for remote water monitoring and detection of coliform.

12 Objectives

The project has the following objectives:

- i. To develop and construct a portable incubator capable of maintaining a specific temperature range of 35-37°C using a ESP32-powered relay to control the heating element and sensor readings.
- ii. To detect coliform bacteria growth using an LDR sensor and a bulb by monitoring the changes in resistance.
- iii. To develop a real-time monitoring and data visualization of coliform levels.
- iv. To validate the accuracy and effectiveness of the portable incubator in detecting coliform bacteria.

1.5 Methodology

The proposed research falls under the field of environmental science and seeks to examine the quality of water samples that are already contaminated with coliform bacteria. The project will utilize the Waterfall model of methodology, an iterative process for creating a system or solving a problem that structures the entire job into segments to ensure precision.

1.5.1 Waterfall model of methodology

This project has been successfully accomplished by implementing the following strategy:

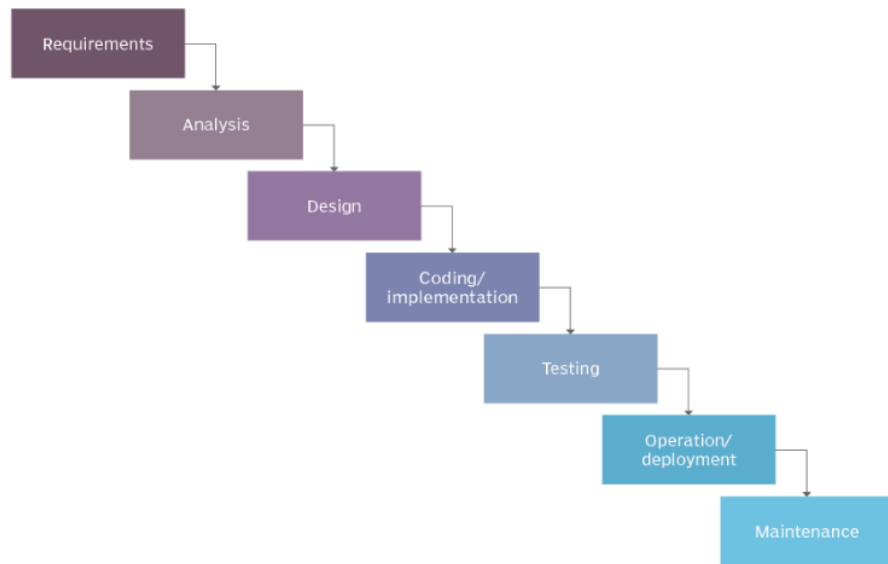


Figure 1.1: Waterfall model

1.5.2 Description of the Waterfall model

The work has been grouped into seven sections;

- 1 **Requirements Gathering Phase:** In this phase, the aim and objectives of the project will be defined, and the specific requirements for the research will be gathered.
- 2 **Analysis Phase:** In this phase, the water samples containing coliform will be analyzed, and the most suitable water samples for the experiment will be identified and selected based on specific criteria.
- 3 **Design Phase:** In this phase, the portable incubator will be designed, and the DHT11 sensor and LDR will be integrated into the system.
- 4 **Implementation Phase:** In this phase, the incubator will be set up, and the selected water samples will be placed in it. The temperature and light exposure within the incubator will be monitored using the DHT11 sensor and LDR, respectively.
- 5 **Testing Phase:** In this phase, any changes in the resistance levels of the LDR due to bacterial growth in the water samples will be recorded.
- 6 **Deployment Phase:** The data acquired during the testing phase will undergo statistical analysis to identify and quantify the existence of coliform bacteria in the water samples. Statistical methods will be employed to assess the presence and concentration of coliform bacteria based on the collected data. 9
- 7 **Maintenance Phase:** During this phase, any required modifications to the experiment will be implemented in response to the discoveries and outcomes of the investigation. Finally, conclusions and suggestions will be drawn based on the results obtained.

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1.6 Scope of Study

The primary focus of this research is to conceive and create an economical and easily transportable incubator tailored for remote monitoring of water quality and identification of coliform bacteria. The incubator will utilize a light-dependent resistor (LDR) to detect changes in turbidity, and a DHT22 sensor to maintain a consistent temperature of 35-37°C inside the incubator. The incubator's operation will rely on a 230 AC – 5V DC power adapter and monitored using ESP32 module. The study will involve designing and constructing the incubator, testing its sensitivity and specificity for detecting turbidity increases, and evaluating its performance in a real-world setting.

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The study aims to provide a cost-effective and efficient solution for monitoring water quality in remote areas where access to laboratory equipment is limited.

1.7 Limitations of the Study

Limitations associated with this project include:

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- i. The precision of coliform detection might be influenced by various factors, including the quality of the water sample and the calibration of the LDR sensor.
- ii. The use of an ESP32 microcontroller for data transmission may have limited range and reliability in remote areas with poor internet connectivity.
- iii. The incubator may require frequent calibration and maintenance to ensure consistent temperature and humidity levels for accurate readings.
- iv. The heating source required to increase the temperature in the incubator uses up a significant amount of power, which makes it impractical to use a battery as a power source.
- v. The size and portability of the incubator may limit the amount of water samples that can be tested at once, potentially affecting the scalability of the system.

1.8 Project Organization

The report is structured into five chapters, each contributing to a comprehensive understanding of the project. Chapter One serves as a broad introduction, providing a general perspective on the research, contextual background, specific goals and objectives, targeted issues, and a concise outline of the chosen methodology. In Chapter Two, a thorough literature review is presented, delving into relevant studies to establish a theoretical framework and provide additional insights crucial for a comprehensive grasp of the project. Chapter Three focuses on the project's methodology, discussing the necessary components, specifications, design considerations, and innovations, including system block diagram representation and hardware design. Moving on to Chapter Four, the implementation and testing of the project are covered, showcasing the practical system design and featuring in-progress photos. This chapter also entails a comprehensive examination and analysis of the project's findings and outcomes. Finally, Chapter Five serves as the concluding chapter, summarizing the study's conclusive remarks, outcomes, and displaying the project's achievements.

CHAPTER TWO

LITERATURE REVIEW

2.1 Introduction

Many researchers have investigated the prevalence of coliform bacteria in various sources of water. Coliform bacteria existence in drinking water is a huge public health concern. Although total coliform bacteria are widespread and typically not harmful, the presence of fecal coliforms and *E. coli* may pose a potential danger to human health. Coliform bacteria, including total coliforms, fecal coliforms, and *E. coli*, are frequently employed as indicators of water quality. [3]. This chapter provides a comprehensive overview of Coliform bacteria, including the major factors that contribute to their presence in water, the potential health effects of exposure to these bacteria, traditional methods for detecting them in water, relevance of IoT technology in the detection of Coliform bacteria in water and review similar works in the literature.

2.2 Coliform Bacteria: A Quick Overview

Coliform bacteria are rod-shaped, non-spore-forming, Gram-negative, oxidase-negative, facultatively anaerobic organisms that have a cell wall appearing pink or red when stained and can survive in both oxygenated and non-oxygenated environments [4]. Coliforms are a class of bacteria that are present in soil, plants, and the digestive tracts of all animals, including humans. While most coliforms are harmless, the presence of certain coliforms, like *E. coli*, in drinking water might signal that the water may be polluted with feces, which increases the risk of illness. The presence of additional pathogens is possible if *E. coli* is discovered in the water, since *E. coli* itself can be a pathogen. The intestine of warm-blooded mammals is where it typically began

[5]. In the past, the presence of the intestinal bacteria known as coliforms has been cited as a sign that a location has been contaminated by sewage from people. Additionally, the feces of affected people contain harmful germs including Shigella and Salmonella[6].

Coliform bacteria are classified into three types that indicate the safety of water and differ in the level of risk. The term "total coliform" encompasses a wide range of bacteria, including those present in fecal matter, which is a subset of all coliforms. One variety of fecal coliform bacteria is E. coli. When testing drinking water samples, laboratories check for the presence of total coliform, and if it is detected, the sample is further examined see whether it contains E. coli[3]. Total coliform bacteria presence is commonly observed in nature and does not pose a significant threat. Drinking water contamination is most likely environmental when only total coliform bacteria are found in it[7].

Total Coliform Bacteria: The common bacteria in the ecosystem known as total coliforms is usually found in dirt or plants, and is widely regarded safe. If a laboratory only observes total coliform bacteria in drinking water research, then it is likely that the bacteria came from the environment and fecal pollution is rare. However, there is a chance that harmful germs could get inside the system if it is prone to external pollution. As a result, It is crucial to identify and address the primary cause of the contamination.

Fecal Coliform Bacteria: In the intestines and feces of both humans and animals, fecal coliform bacteria, a subtype of total coliform bacteria, are prevalent. The identification of fecal coliforms in a water sample is typically indicative of recent fecal

contamination, and this poses a higher risk of potential pathogen presence compared to situations where just total coliform bacteria are identified[7].

Escherichia coli: The bulk of E. coli types usually pose little worry and typically live [35] in the intestines of warm-blooded creatures like humans. E. coli is a subset of fecal coliform germs. However, some kinds of E. coli can make people ill. When E. coli bacteria are found in a portable water sample, it usually means that there has been recent fecal pollution, which boosts the chance that disease-causing germs are present [84] [7]. The identification of E. coli in water samples typically indicates contamination with feces, along with the potential existence of human-originating harmful organisms [7]. Since E. coli can also be a pathogen, there is a possibility that other pathogens will also be present if E. coli is discovered in the water[5]. Escherichia coli bacteria were typically found in sewage discharges from human, animal, and plant sources, in that order. Septic system failure, municipal landfills, and wastewater sludge application to the land are all examples of human sources. Numerous animal sources, including domestic pets, wild animals, cattle, the spreading of manure on land, pastures, and feedlots, can release fecal coliform bacteria into the environment[8].

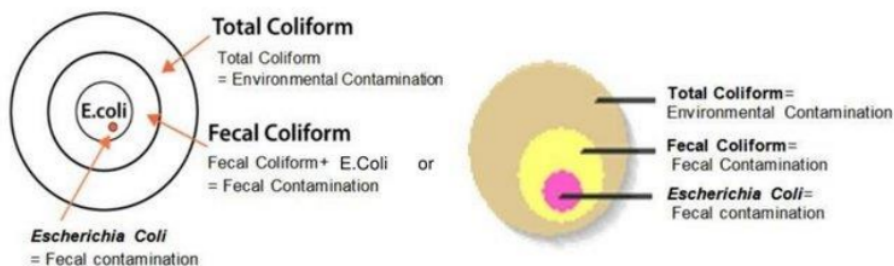


Figure 2.1: Different Groups of Coliform Bacteria [7]

2.3 Significant Factors that promote Coliform Bacteria infestation in water

Water sources containing coliform bacteria pose a significant risk to the public's health and are frequently contaminated by feces due to improper sanitation practices or agricultural activities. It has been claimed that septic tank systems and farm waste are the main contributors to coliform bacteria in drinking water. Septic tank overflow and other related factors contaminating drinking water sources with human feces particularly raise the risk of Coliform contamination. The risk of contracting a waterborne disease can also be increased by the use of pesticides and fertilizers in agriculture, which can reduce the spread of coliform bacteria to water sources. To protect public health and stop the spread of infectious diseases, coliform bacteria must be found and monitored in drinking water.

2.3.1 Septic Tank Systems.

20 Water contamination is one of the biggest problems in the world today. Septic tank effluent intrusion is one of the main causes of groundwater contamination. the principal origin of harmful viruses and bacteria in the subterranean environment may be septic tank effluent[9]. Contamination of aquifers has become a common problem as a result of excessive groundwater pumping and uncontrolled human wastewater dumping into the groundwater. Groundwater quality can be impacted by residential, business, industrial, and agricultural activities. Groundwater is particularly sensitive in regions with large population densities and extensive human use of the land[10]. Waste is temporarily stored in a septic tank while being pre-treated at that time.

There have been reports of septic tanks failing and leaking a lot, harming the ecosystem[11]. Researchers from several fields have done studies to show the detrimental effects of situating septic tank systems close to drinking water sources. An examination 23 investigating the impacts of septic tank distances from wells and groundwater

contamination in the Agbowo suburb of Ibadan, Nigeria, was done[12]. Septic tanks have been proven to malfunction and leak a lot, harming the environment [11]. Therefore, septic tanks are the primary source of waste that is directly released into groundwater. Monitoring of water is therefore crucial to protect both the environment and human health[13]. Forty (40) ground water samples were replicated, and they were taken from groundwater sources at different distances for bacteriological analysis, and it was found that they were all infected with different types of bacteria as well as coliform[12]. The number of feces and total coliforms decreased with increasing distance from septic tanks, and this effect was more pronounced for water supply distance from pollution sources. Because the problem of septic tank contamination of ground water is a national concern rather than a local one in the study region, Standards ⁶⁵ should be created for the separation of wells from septic tanks and the pretreatment of well water[12]. Consuming groundwater that has been faecally contaminated and not properly treated has been related to disease[14].

⁷²Underground water, or the water found below the earth's surface in soil pores and rock formation fractures, has been impacted by the degradation of water's physicochemical and biological properties brought on by industrialization and urbanization, which have grown over time without consideration for the effects on the environment[15]. As a result of population expansion and urbanization, ground and surface water are subject to greater demands. The startling rate of urbanization in Nigeria is reflected in the 10–15% yearly growth rates of the country's main cities[16] hence human activities like enhancing soil fertility, irresponsibly disposing waste, and utilizing pit latrines, soak-away pits, and septic tanks are increasing. In Nigeria, pipe-borne water is hard to find, thus a lot of homes use wells that are close by but separate from septic tank. Adequate drinking water sources are accessible to only 52% of the population in

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Nigeria[17]. The proximity of septic tanks to wells and the unsanitary conditions in the area around the wells are the main causes of contamination. Some wells are poorly maintained and lack sufficient coverings or lids, making the water unfit for consumption and raising the possibility of waterborne infections, and the unhygienic use of the wells. Additionally, the discharge of solid or liquid waste into pits, stream channels, abandoned boreholes, landfills, or other unlined sites may result in groundwater contamination[18].

The authors of a different study investigated the origins of E. coli contamination in the coastal subtropical climate. They found that septic systems were a substantial

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source of E. coli and total coliforms in both surface water and sediment. It was evident that septic systems were a substantial source of pollution since there was a
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positive correlation between the amount of E. coli in sediment and the distance from them. The authors asserted that septic systems are a significant source of fecal
25
contamination in coastal areas and suggested introducing measures to guard against contamination from septic systems[19].

Another study looked at how the presence of septic tanks affects the quality of the borehole water in Port Harcourt, Nigeria[20]. In order to reach their conclusions, the researchers collected water samples from boreholes both close to and distant from
46
septic tanks. Total dissolved solids (TDS), pH, total suspended solids (TSS), and bacterial contamination were all investigated in the water samples (TSS). The researchers found that the water quality in boreholes near sewage tanks was significantly lower than in boreholes farther away from septic tanks. The water samples collected from the vicinity of septic tanks contained higher concentrations of TDS, TSS, and bacterial contamination, including coliform bacteria[20].

Another study used one-way ANOVA (Analysis of Variance) looked into the connection between fecal coliform contamination and proximity to the septic tank. The findings revealed that 96 percent of the examined wells had fecal coliform, indicating that the drilled well's biological drinking water quality had been significantly diminished. The excavated well was 53.7 feet away on average from the septic tank.

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The results of the investigation showed that as the distance between the septic tank and the drilled well rose, fecal coliform contamination level dropped. There were strong evidence of fecal pollution in the dug-well water at 50% of the septic tanks within 50 feet[21].

1. Septic tank setback distances from water sources

Since the distance between septic tanks and water sources plays an important role in coliform bacteria which causes water to become contaminated, various approaches for estimating setback distances and conditional probabilities have been examined. To place restrictions on the permitted places for septic tank placement, government agencies have implemented laws stipulating minimum setback lengths between septic tanks and drinking-water wells[22]. How likely is it that a setback distance (such one required by the regulatory body) will be adequate to protect the city's ground water from virus contamination? Given a specific probability level, what setback distance would be necessary to ensure that the ground water would be immune to virus contamination?[23] With little respect for the local geology, hydrology, and meteorology, these setback distances are often required for at least a county-wide area[23]. Various places have different topologies. Therefore, proper setback distance calculations must be made by regulatory agencies to be able to guarantee that the distance of septic tanks from water sources would give complete assurance that the water is not contaminated with coliform. A study revealed that a setback distance of 40

m would be required for a specific location under consideration in order to be 90% confident that the ground water would be appropriately protected from virus contamination. If one wanted to be 99% sure that the setback was enough to stop viral transmission in places where 15-m distances were first permitted, an 80-m distance would be required[23].

2. Inadequate agriculture practices

Numerous studies have demonstrated how agricultural pollution may significantly contribute to the coliform water contamination. One of the main sources of contamination is the over use of fertilizers, notably those containing nitrogen and phosphorus. According to the National Geographic Society, nutrient contamination brought on by excessive fertilizer use leads to eutrophication, a process that harms both terrestrial and aquatic ecosystems by overburdening them with nutrients[24]. Coliform water contamination is a global problem that threatens the health of both people and animals. Ineffective agricultural techniques have been identified as one of the main sources of this problem. For instance, it is common knowledge that increasing soil fertility with animal manure also increases the quantity of fecal coliform bacteria in the water. Therefore, excessive fecal coliform bacteria levels and other pathogens can contaminate surface and groundwater when animal dung is improperly handled or applied[25]. Surface water, such as streams and rivers, can be contaminated by runoff from feedlots and farms, and groundwater can be affected by pollutants that seep through the soil layers. People and animals may be exposed to toxins while using contaminated groundwater for irrigation or drinking. Different management techniques can be used to reduce the amount of water pollution caused by agricultural activities. Among these include reduced use of pesticides and fertilizers, conservation tillage, nutrient management programs, and manure management strategies. Additionally, by

creating buffer strips or wetlands or increasing irrigation effectiveness, runoff can be reduced and contaminants in surface water can be filtered[26]. Water sources may be impacted by pesticides, herbicides, and fungicides used in agriculture. It is well recognized that these agricultural contaminants deteriorate groundwater and surface water quality[27]. Using contaminated irrigation water and disposing of animal corpses inappropriately are two other poor farming techniques that could cause coliform contamination[28]. Rainfall and flooding are other causes of water contamination. They accomplish this by introducing potentially harmful chemicals and contaminants into water sources from a variety of sources. Studies conducted in the lab and in the field indicate that there are significant numbers of bacteria in water, which are primarily brought on by frequent tidal floods and strong rain events[29]. Inadequate water consumption spurred on by population increase, urban wastewater runoff, hospital waste, and institutional waste are other causes of coliform contamination of water[21].

Poor hygiene habits, such as defecating outside, and poorly maintained wells that have free access make them more susceptible to anthropogenic pollutants[30]

2.4 The significance of finding Coliform bacteria in water

Monitoring is the primary line of defense against diseases brought on by dangerous microbes. Effective detection techniques are required to manage this effect[31].

To determine whether there are any pollutants in the water, accurate and quick reaction detection technologies are required[32]. Water safety and cleanliness are basic human rights and needs for all living beings[33]. The three most basic uses of water—drinking, food preparation, and personal hygiene—require each person at home to use 7.5 liters of water per day at least. A minimum water consumption of 50 liters per person per

day is necessary to meet the requirements for personal hygiene, food hygiene, housekeeping, and laundry purposes[34]. However, due to its importance as a way of propagating pathogenic organisms, man is now worried and terrified of it[35]. Animal and human waste is a significant contributor to water pollution, which is extremely hazardous to the health of millions of people worldwide. Waterborne illnesses like cholera, polio, diarrhea, hepatitis, typhoid, dysentery, and gastroenteritis are brought on by these microorganisms that contaminate the water supply[36]–[38]. One of the top 15 global killers is diarrheal disease[39]. More children die from diarrhea than from tuberculosis and malaria combined[34]. Millions of people globally are afflicted by other ailments linked to WASH. Over 800,000 deaths are attributed to inadequate WASH each year, particularly among children under five[40]. Elderly and young people often suffer more from the harmful health effects of drinking water contaminated with *E. coli*[41]. The difficult work of assessing and analyzing the microbiological features of water in rural developing areas is one of the major barriers to promoting and implementing safe water projects. Many remote populations lack the necessary tools and knowledge to evaluate the microbiological purity of water before using it. Despite the large number of hazardous microbes, testing for all of them is challenging[42]. To reduce dependence on laboratory testing and overcome the challenges of cost, logistics, and time delays associated with sending samples offsite for analysis, especially in remote areas lacking reliable access to electricity or nearby laboratories, there is a strong demand for simpler, faster, and cost-effective methods that enable frequent water testing at the community or household level.[43]

2.5 Traditional methods of detecting Coliform in water

Drinking water is an oligotrophic system, thus the inability of culture methods to recognize starved and stressed bacterial cells could have serious repercussions by drastically underestimating contamination levels. There are other alternative methods for recognizing coliforms, albeit they are all at different stages of development and implementation. In this overview, cutting-edge and new methodologies are discussed alongside the fundamental principles and accepted practices of traditional methods of Coliform detection. Additionally, based on each method's benefits and drawbacks, the efficacy of finding coliforms in oligotrophic environments, such drinking water, is assessed. Additionally, criteria like detection sensitivity and limit, amount of time needed to collect data, and laboratory costs are taken into account (including skill, labor, and cost)[4].

Traditional methods include membrane filter method, multiple-tube fermentation method, enzymatic/defined substrate method and molecular methods.

2.5.1 Membrane filter method

The membrane filter method is one way for calculating the quantity of microorganisms in drinking water which has achieved acceptance in a number of countries. Membrane filter technology is used to keep an eye on water quality. This method comprises running a water sample through a 0.45 mm-pore filter that is sterile and bacteria-retentive. After the filter has been cultivated on a selective medium, the typical colonies on the filter are then counted. To find the optimum method for recovering coliforms from water samples, various media and incubation conditions were examined. The Tergitol-TTC medium and the m-Endo-type media, respectively, are frequently used for drinking water analysis in North America and Europe[44]. After 24 to 48 hours of incubation, coliform bacteria form yellow-orange colonies on Tergitol-TTC media,

whereas they create metallic-looking red colonies on an Endo-type medium that contains lactose. While studies indicate that m-Endo agar is associated with greater efficacy in cultivating higher quantities of coliforms, alternative media like MacConkey agar and Teepol medium have been utilized in South Africa and the United Kingdom. In the determination of fecal coliform counts, filters are commonly incubated for 24 hours at 44.5°C on an enhanced lactose medium known as m-FC[44].

2.5.2 Multiple-tube fermentation technique

Over the past 80 years, the multiple-tube fermentation (MTF) approach has become a popular methodology for enumerating coliforms and monitoring water quality. In this method, multiple test tubes containing various dilutions of the water sample are employed. The occurrence of gas generation, acid formation, or noticeable growth in the test tubes after a 48-hour incubation period at 35°C indicates a positive presumptive reaction. Both lauryl tryptose broths and lactose broths can serve as presumptive media. Confirmation tests are conducted on all tubes once a positive presumptive result is observed. To pass a confirmation test, a brilliant green fermentation tube containing lactose-bile broth must exhibit gas production within 48 hours at 35°C[4].

The multiple tube fermentation method outcomes are expressed using the most probable number (MPN), which is a statistical estimation of the mean number of coliforms. Using this technique, it is possible to determine the coliforms semi-quantitatively. The measurement's level of precision is poor because few tubes were used for the analysis. The likelihood of getting a negative result when using five tubes, each with a volume of 1 ml of the material, is less than 1%.

Despite its limitations, the MTF technique is still widely used and a good way to check water quality. Some of its benefits include simplicity and low expense. However,

compared to the defined substrate approach, this technique has a significant disadvantage due to its lengthy incubation period. The MTF technique is a vital instrument for ensuring the safety of drinking water because it can identify both total and fecal coliforms. However, the MTF technique's lack of accuracy, brought on by its reliance on statistical estimates, raises the possibility that it may not be the best technique in circumstances where high precision is required. Alternative approaches with higher accuracy, like PCR-based techniques, may be preferred in such cases.[4]

2.5.3 Molecular methods

There are currently several molecular techniques available that enable rapid, sensitive, and precise identification of pathogenic microorganisms in clinical samples and assessment of the microbiological safety of food and water. These methods have been developed and refined over the past two decades. Unlike traditional approaches that involve isolation and cultivation of microorganisms, these molecular techniques allow for qualitative and quantitative analysis of the target microorganism without the need for growth. Moreover, these methods can be customized to detect specific pathogenic traits carried by the microorganism. Various target molecules can be utilized, and the ultimate objective of these techniques is to provide valuable insights for the prevention of infectious diseases[45]. The following are the molecular techniques used to find coliforms in drinking water

1. Polymerase chain reaction method

The polymerase chain reaction (PCR), a method that is frequently used to make millions to billions of copies (full or partial) of a given DNA sample, allows researchers to quickly amplify a very small DNA sample (or a portion of it) to a huge enough quantity to analyze in depth.[46]. An amplified DNA target segment

can now cycle through replication thanks to this reaction. Oligonucleotide primers are used for replication in a chain reaction that is started by a DNA polymerase, specifically Taq polymerase. Both in vitro and in situ replication methods are possible. The degree of complementarity and homology between the target sequence and the primer, as well as the temperature at which hybridization occurs, affect the sensitivity of the detection method.[44]. By using gene probe detection and PCR amplification of particular portions from two genes, lacZ and lamB, the identification of coliform bacteria was investigated. *Salmonella* spp. and non-coliform bacteria were not found when *Escherichia coli*'s lacZ coding area was amplified using PCR primers with an annealing temperature of 50 degrees C. However, this technique was successful in detecting *Shigella* spp. as well as other coliform bacteria. The identification of *Salmonella*, *Shigella*, and *E. coli* species was accomplished by selectively amplifying a segment of *E. coli* lamB at an annealing temperature of 50 degrees C[47].

2. Nucleic Acids

For accurate identification of certain phylogenetic sources and species identification, DNA and RNA, particularly nucleic acid polymers, are very suitable molecular targets. Numerous molecular techniques for coliform detection have been developed thanks to the stable genotype properties of nucleic acids[45]. Coliforms in drinking water must be found in order to protect public health, and nucleic acid-based molecular methods has established itself to be a quick, sensitive, and reliable way to do so[44]. Overall, nucleic acid-based molecular approaches provide a number of advantages over traditional culturing methods in order to recognize coliforms in drinking water. They don't require labor- or time-intensive culturing methods and can identify coliforms rapidly and precisely. These techniques are essential because they reliably identify the

presence and amount of coliforms in water samples, protecting public health, and preventing the spread of infectious diseases [44].

2.5.4 Enzymatic method/defined substrate method.

The Colilert-18 is one of the most often used defined substrates for the enzymatic test because it contains the nutritional indicators ONPG and MUG from Defined Substrate Technology (DST), which establishes *E. coli* and Coliform bacteria presence. The US EPA has authorized the reagent as a routine procedure for evaluating treated and tainted water since it may deliver results in just 18 hours. With the use of this reagent, you may attain 100% test accuracy. Colilert-18 reagent will be entirely soluble in water after being shaken into water samples[48]. Other efficient substrates, such those created by the Italian business Liofilchem, are already on the market and are good indicators of coliform in water. Incubation of samples is done for 18–24 hours at 37°C or close to 48 hours at room temperature (20–25°C)[49]. Coliforms have the enzyme β -galactosidase, which they use to create gases and acids when the temperature is between 35 and 37 °C, which is ideal for growth[50]. A certain enzyme will produce color or fluorescence when it digests a chromogenic or fluorogenic substrate. These substrates have been used to determine whether or not certain enzymes are present or active in water. To detect the existence of b-D-glucuronidase in *E. coli*, scientists employed a chromogenic substrate called indoxyl-b-D-glucuronide (IBDG). Various commercial tests based on this specific substrate technology were subsequently created, such as Colilert (IDEXX Laboratories, Portland, ME, USA), Colisure (Millipore Corporation, Bedford, MA, USA), and Coli-Quick (IDEXX Laboratories, Portland, ME, USA) (Hach, Loveland, CO, USA)[4].

2.6 Internet of Things (IOT): A Quick Overview

The phrase "Internet of Things" (IoT) refers to a network of physical things that are integrated with sensors, software, and other technologies, enabling them to connect and communicate with other devices and systems via the internet without requiring human intervention. These devices cover a wide range of products, from standard domestic appliances to cutting-edge industrial machinery[51]. These objects have the ability to perceive, communicate, interact, and function together without requiring human-to-human or human-to-computer interaction thanks to integrated technology and a network of physical objects[52]. IoT is about more than just connected devices; it's also about the data those devices gather and the immediate, potent insights that can be drawn from it [53]. Internet of Things (IOT) increases productivity since it reduces reliance on humans[54]. This program has been used to develop programs for monitoring ranging from surveillance, healthcare, and the environment all around the world[55].

2.6.1 Structure of the Internet of Things (IOT)

Sensors, microcontrollers, and application platforms make up the IOT architecture.

1. Sensors

IoT sensors track environmental changes in real time by detecting the physical environment[56]. IoT sensor networks' main objectives cover three areas: (i) gathering important data from the outside physical world; (ii) sampling internal system signals; and (iii) using sensor data to deliver appropriate information for decision-making[56], [57]. These sensors have nodes built in that are linked to microcontrollers that are in charge of reading and analyzing the data collected by the sensors. The most often used IoT sensors are those that measure temperature, pressure, humidity, level, accelerometer, gas, gyroscopes, motion, picture, and optical properties, as well as RFID (Radio Frequency Identification) and infrared (IR) sensors[56]. The small size of the

sensor causes technical restrictions in areas like computational power, networking ability, battery life, storage capacity, and memory, despite the fact that sensors have a wide range of applications. This means that these sensors could be very vulnerable to errors, assaults, and simple failure, which could lead to the loss of sensor data and erroneous information[56].

2. Microcontroller

A microcontroller is a small computer with network access and program execution capabilities. The controller manages the collection of numerical data from the sensors, which it can then upload to the cloud or the internet for storage and analysis[58], [59]. Minicomputer- and microcontroller-board-based IoT controllers can be distinguished from one another. Microcontrollers like Arduino are more affordable than minicomputers and can run the same program repeatedly. The ESP is an additional instance of a microcontroller (ESP32, ESP8266). As an alternative, portable, full-featured computing devices that are capable of handling a variety of tasks include minicomputer-type controllers like the Raspberry Pi. While an Arduino board requires additional external hardware in order to link up with the internet, the Raspberry Pi comes with an Ethernet and Wi-Fi network interface built in[59].

3. Application Platform

Without applications, IOT is of no meaning. Applications for Internet of Things provide reliable communication and real-time message transmission. An application is essentially a piece of software that runs online and, after receiving input from a microcontroller, displays data via a user interface. Platforms such as Blynk offer mobile applications that enable the management and monitoring of IoT devices via mobile devices. Additionally, FreeBoard, Ubidots, and ThingSpeak are three other well-known platforms utilized for IoT applications [59].

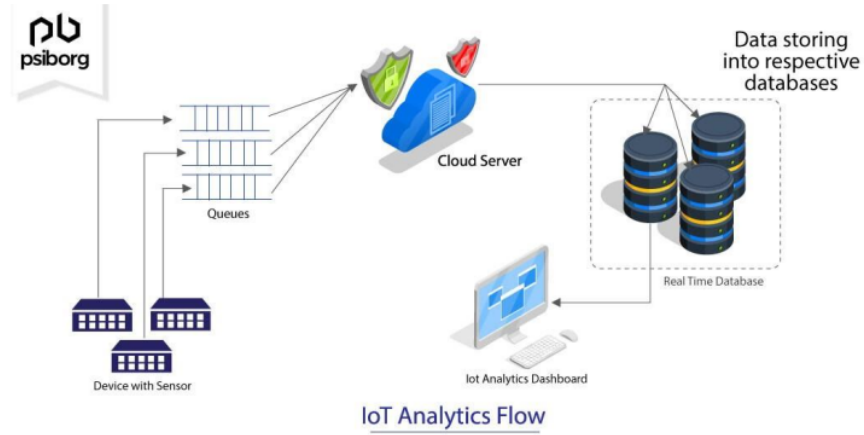


Figure 2.2: IoT Analytics Flow [60]

2.6.2 IOT's relevance for detecting bacteria in water and its utility in detecting coliform

Water samples and the data that go with them are often processed and examined in laboratories. This may take a lot of time and effort and result in less trustworthy results.

In light of the situation, it is essential to establish a suitable and organized system for continuous monitoring and management of water resources in order to address connected challenges. Internet of Things (IoT) enables the collection of real-time remote measurements which may be conducted with little to no human participation, providing a great alternative to conventional, complicated, and unsuccessful techniques for reaching this goal[61]. Due to the harmful health implications of consuming contaminated water, real-time monitoring of water quality is now required. IoT domain addresses this problem by providing creative and reasonably priced solutions for the problems of water contamination detection and water quality analysis[62].

The following are some benefits of using IOT for water quality monitoring:

- Decreased cost of computing:** The considerable amount of computing work, like that required by the Ubidots IoT-Platform, is handled by cloud servers.

With the passage of this law, any widely used processor-based node (such as the NodeMCU (ESP32) kit) may be used in place of sophisticated microprocessors[63].

2. **Hardly any human involvement:** IOT systems can increase efficiency by reducing reliance on humans[64].
3. **Low cost:** IoT solutions are generally more cost-effective than traditional water monitoring techniques since they make use of currently in use, commercially available communication infrastructure, which is also more time-consuming[63].
4. **Low power requirements:** The local nodes only need to carry out minimal work after shifting the majority of the computing workload to the cloud server, and they can then primarily sleep. This made it possible to operate sensor nodes in far-off places with a little battery.
5. **Real-time feedback delivery:** The proper water authority and end users can both immediately receive real-time input from these systems via SMS, email, Twitter, Facebook, and other means[63].

2.7 Review of related works

This section analyzes experiments that involved the development of an improvised incubator for coliform growth in contaminated water samples as well as the identification of coliform and other types of bacteria in water using Internet of Things (IoT) technology. The next paragraphs concentrate on a few of the numerous research studies that have been carried out in this field.

A sensor-based Water Quality Monitoring System was employed in another study[52] to measure the water's molecular and physical properties, including its pH, temperature,70

turbidity, and Total Dissolved Solids. (TDS). The authors' IoT-based solution consisted
of three crucial components: a multiparameter water quality analyzer, a means for
information transmission, and a computer system unit. Sensors, Wi-Fi wireless
transmission, an STM32 single chip microcontroller, and remote water quality
monitoring were all included in the proposed system. The data on water quality
collected were evaluated using ANOVA-based statistical analysis to identify the key
causes of the river Krishna's deteriorating water quality. The authors used a water
quality monitoring system based on the Internet of Things (IoT) to gather and evaluate
current data on the river Krishna's water state in order to identify the primary causes of
water contamination.

IoT-based technique for predicting bacterial contamination in water was proposed by
the authors of a study paper[61]. The setup entails placing a variety of equipment in
the study area, including sensors for determining total dissolved solids, pH level,
turbidity, and dissolved oxygen. These tools are used to evaluate and keep track of the
water's quality. These sensors transmit the data they have collected to a server using
GSM modules, where it is processed and examined. The scientists used machine
learning methods to predict coliform bacteria presence in the water based on the sensor
data collected by the IoT system. The proposed approach provides a more useful and
cost-effective alternative to manual lab analysis, which is an expensive and time-
consuming process for the prediction of bacteria. To predict coliform presence bacteria
in water, the authors created an IoT-based system that uses a variety of sensors and
machine learning techniques, providing a quick and cheap alternative to traditional lab
examination.

An incubator powered by batteries that effectively sustains a temperature range of
approximately 35°C with a tolerance of $\pm 1^\circ\text{C}$ of 25°C for up to 65 hours is described

in another study[65] free, open-source design. The objective of this incubator is to provide humanitarian engineers with a quick, simple, and visually pleasing method to evaluate the quality of microbes in water in remote locations. To confirm the presence of E. coli in the water samples, the authors used 3M Petrifilm E. coli/Coliform (EC) Count Plates. These dishes need to be incubated at a temperature that is in close proximity to body temperature. In order to meet this need, the portable incubator was developed, and the authors demonstrate that it can maintain the required temperature range for up to 65 hours. However, the use of IoT in the research is not acknowledged by the authors. As has been shown, IoT has the ability to improve water quality monitoring in various ways. Using temperature, pH level, dissolved oxygen, and turbidity as an example, Internet of Things (IoT) sensors can provide real-time information about these factors that affect water quality. This data may be sent to a database for adequate analysis and visualization, allowing for swift and educated decision-making. Additionally, IoT can allow remote water quality monitoring, doing away with the need for labor-intensive manual analysis.

In a different study, the writer used IoT to monitor and track the water quality in real-time[58]. They suggested a system that uses IoT sensors to measure pH, temperature, and turbidity in addition to other elements of water quality for use in homes. The system's objective is to offer remote internet access for managing and evaluating water purity, for instance through a mobile application. The proposed system uses Big Data Analytics and Machine Learning techniques to analyze water quality data in-depth, allowing the early detection of threats. In conclusion, the research's authors used IoT sensors and big data analytics techniques to measure and track water quality in real time, making it possible to identify risks early enough to prevent them and to manage water quality remotely.

In a different study[48], the authors developed a portable incubator for the development of coliform and E. coli germs using IoT technology. With the aid of this portable incubator, microbiologists can evaluate the quality of the water anywhere—not just in laboratories. Using qualitative and quantitative techniques, the incubator is able to identify E. coli in water samples as well as coliform bacteria. Two-factor authentication security enables real-time tracking, and the incubator can cut the detection period to just two hours. (2FA). The capability of the device to reach and maintain a predetermined temperature (typically 35–37°C) under various conditions, a procedure known as bacterial growth, determines the device's dependability. The incubation procedure can be carried out manually or automatically via Internet of Things (IoT) technology and computer vision. Data is protected by encryption techniques in the incubator, and the connection between the device and Wi-Fi is also secure. The outcomes of their experiment can also be observed in real time using IOT apps. In conclusion, the researchers developed a movable E. coli and coliform bacteria incubator using IoT technology that can identify the presence of these bacteria in water samples using both qualitative and quantitative techniques. Because it is dependable, secure, and can cut the detection time to just two hours, the incubator is an essential tool for microbiologists to use when conducting water quality checks all over the world.

In a different study[66], the authors describe a method for building a movable, affordable incubator for drinking water microbiological testing. The incubator prototype can be modified to handle different materials as well as volumes and has a disassemblable design that makes carrying it easy. This may be used in grid-powered, laboratories as well as remote areas with solar or automobile batteries. The design uses components that are typically available, can work in a variety of outdoor settings, and maintains the advantages of more expensive laboratory-based models. In order to

assess the microbes in drinking water, the writers wished to address the need for a portable, affordable incubator. A variety of culture-based microbiological techniques, 29 including membrane filtration and culturing, are used to determine if water is fit for human consumption, can be used in the incubator. Overall, the authors' work suggests that their design may provide a practical and affordable alternative for the incubation of microbial colonies in the lab and the field. Their approach has the potential to be extensively applied in low-resource areas where it may be difficult to obtain specialized equipment. However, they did not incorporate any kind of IOT into their architecture.

The authors of another study [67] proposed a low-cost light detecting unit-based turbidity instrument for monitoring water cloudiness. The system makes use of an Intel 44 Galileo 2 processor and a server for web-based tracking. The Turbidity Detection Unit 31 is made up of a polyvinyl chloride (PVC) conduit, and a Light Emitting Diode (LED), a Light Dependent Resistor (LDR). The turbidity system relies on the idea that the amount of suspended matter and the amount of light it dissipates are directly correlated

51 The authors propose a clever Internet of Things (IoT)-based system for monitoring water quality indicators in another research publication [68]. The device is designed to monitor four physical water quality parameters: Turbidity, temperature, electric conductivity, pH. An Arduino microcontroller, Sensors, an ESP8266 Wi-Fi module, and a cloud database for data storage and analysis make up the proposed system. The device operates by using sensors to collect data from the water sample and transmit it to the Arduino. Using the ESP Wi-Fi module, data is analyzed by the microcontroller and sent to a cloud database. To furnish valuable insights into water quality, the collected data is saved and examined in the cloud database. The recommended system 56 is effective and economical and can be used for real-time monitoring of water quality.

The overall suggestion of the research paper is to put in place a smart water quality

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monitoring system that implements IoT to constantly track important water quality indicators. This technique can help reduce potential contamination exposure while ensuring the availability of clean potable water sources.

CHAPTER THREE

SYSTEM ANALYSIS AND DESIGN

49

3.1 Introduction

This section provides a comprehensive review of the materials, tools, and designs utilized to complete the development of the portable incubator. It offers a detailed explanation of the project's programming, circuit design, and construction methods, including a discussion of the components chosen and the reasoning behind those choices. The chapter also showcases the project's flowchart and algorithm and includes information on the methodology and calculation employed during the building process.

3.2 Components

This project employs three categories of components: electrical, mechanical, and chemical. The electrical components encompass a Light Dependent Resistor (LDR), a DHT22 temperature and humidity sensor, a relay, an ESP32 microcontroller, an LED bulb, a 220-12V (AC-DC) power adapter, a buck converter, a Liquid Crystal Display (LCD), and a potentiometer. The mechanical components consist of the incubator and the beakers utilized for holding the water samples under investigation. The chemical component pertains to the substrate named water detect coliform.

3.2.1 ESP 32

The ESP 32 is an affordable microcontroller that has Wi-Fi and dual-mode Bluetooth built-in. It acts as both the microcontroller and Wi-Fi module, controlling the relay switch and interfacing with sensors. When connected to a Local Area Network (LAN), it enables real-time access to readings through the Blynk platform.



Figure 3.1: Expressif Systems Processor (ESP) 32

47

3.2.2 Light Dependent Resistor (LDR)

This is also known as a photo resistor, photocell, or photoconductor, is a sensor that responds to light changes by altering its resistance. When light falls on an LDR, its resistance changes. The light intensity reaching the LDR is inversely proportional to its resistance. The LDR is used to detect the turbidity levels of the water sample being tested. Due to the water sample being clear initially, more light reaches the LDR, causing a decrease in its resistance. However, if the water sample contains coliform bacteria, after approximately 16 hours of incubation, it turns blue. As a result, the LDR receives less light, causing an increase in its resistance value.



Figure 3.2: Light Dependent Resistor (LDR)

3.2.3 Relay

A relay is an electronic device that can be switched on or off when an external voltage (AC or DC) is applied across its control terminals. In this project, the relay is

utilized to control the heating pad, which helps maintain the desired temperature range of 35-37°C inside the incubator. The heating pad is connected to the relay, which is in turn connected to the ESP32. The ESP32 turns the relay on or off based on the temperature readings obtained from the DHT22 sensor, thereby regulating the temperature inside the incubator.



Figure 3.3: Relay

3.2.4. DC-DC Buck Converter

A DC-DC converter is an electronic device that can increase or decrease voltage of a direct current power source to match a desired voltage level. In this project, a 5V buck converter is used to step down the voltage from the 220V AC to 12V DC power adapter to the 5V level needed to power the various components of the system.



Figure 3.4: DC-DC Buck Converter

3.2.5 PTC Electrical Heating Element

The incubator utilizes an electrical heating pad as its source of heat, which is controlled by the relay. The relay is triggered on and off by the microcontroller, depending on the temperature readings from the DHT22 sensor. The heating pad plays a crucial role in maintaining the temperature range required for the coliform bacteria's optimal growth in the water samples, which is between 35-37°C.



Figure 3.5: Heating Source

3.2.6 LED Bulb

The LED bulb is used as the lighting source. It supplies the LDR with the sufficient light required to determine the turbidity levels of the water samples.



Figure 3.6: LED Bulb

3.2.7 DHT11 Temperature and Humidity Sensor

The DHT11 sensor is utilized to obtain data on the temperature and humidity levels within the incubator. Its significance lies in ensuring that the temperature remains within the 35-37°C range. The ESP32 is connected to this sensor for interfacing purposes.

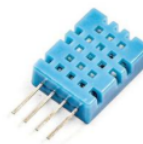


Figure 3.7: DHT 11

3.2.8 Liquid Crystal Display (LCD)

For this project, a 20x4 (20 characters, 4 rows) LCD display is utilized to provide a user-friendly interface for monitoring real-time data readings from various sensors. The LCD display effectively shows temperature and humidity levels within the incubator, as well as the resistance readings of the Light Dependent Resistors. It is interfaced with the ESP32 microcontroller to display the data readings on the screen. The real-time data presented on the LCD enables users to monitor the progress of the incubation process and ensure the optimal conditions necessary for the growth of coliform bacteria.



Figure 3.8:Liquid Crystal Display

3.2.9 I2C Serial Interface Adapter Module

The LCD 20x4 display used in this project is connected to the ESP32 microcontroller using I2C communication protocol. This allows for a simpler and more efficient data transfer between the microcontroller and the LCD display. The I2C interface reduces the number of pins required for communication, making it easier to interface with other components in the project.

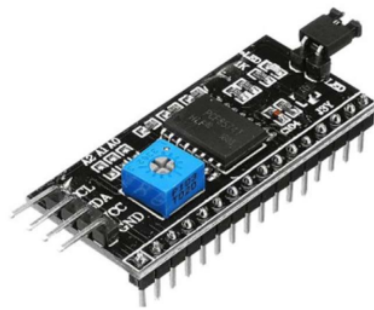


Figure 3.9: I2C Serial Interface Adapter Module

3.2.10 Incubator

The incubator is an essential tool in detecting Coliform bacteria in water samples. Its importance lies in its ability to provide a controlled environment that promotes the growth of Coliform bacteria, necessary for their detection. The incubator ensures accurate and reliable results by maintaining constant temperature and humidity conditions, reducing the risk of false negatives. Additionally, the use of an incubator provides a standardized procedure for Coliform detection in water, allowing for consistent comparison of results across various samples and testing centers. In summary, the incubator is a critical component in the detection of Coliform in water, offering a dependable and consistent method for bacteria growth and detection.



Figure 3.10: Incubator

3.2.11 Beaker

The beakers are utilized to contain the water samples for testing. They are transparent and enable light to pass through the water sample to the LDR. Additionally, they have markings that allow for accurate measurement of the volume of water being tested in milliliters (ml).



Figure 3.11: Beaker

3.2.12 Water Detect Coliform Powder

The chemical being used for detecting coliform bacteria in water samples is a crucial component of our project. It is the only substance capable of detecting the presence of coliforms, which generate acids and gases through β -galactosidase enzyme activity at their optimal growth temperature of 35-to-37-degree Celsius($^{\circ}\text{C}$). The chemical works by breaking down a chromogenic or fluorogenic substrate, resulting in the production of color or fluorescence. Specifically, the powder changes the color of

water samples from a clear and transparent appearance to a dark greenish hue, indicating the presence of coliforms. This color change is important for our project as it allows the Light Detecting Resistors to detect a decrease in light intensity when the water changes color, which indicates the presence of coliforms in the sample.



Figure 3.12: Water Detect Coliform Powder

3.2.13 Power Supply Adapter

The incubator's components are powered by a power supply adapter, which includes a rectifying circuit that converts a 220V AC power supply to a 12V DC power supply suitable for our components.



Figure 3.13: Power Supply Adapter

3.3 Design specification

3.3.1 Systems Block Diagram

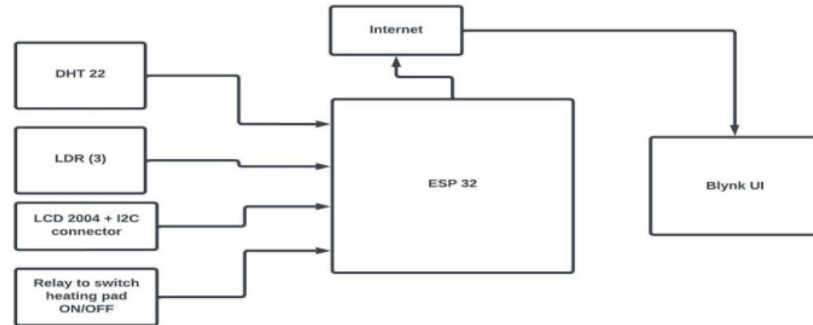


Figure 3.14: Systems Block Diagram

3.3.2 Circuit Diagram

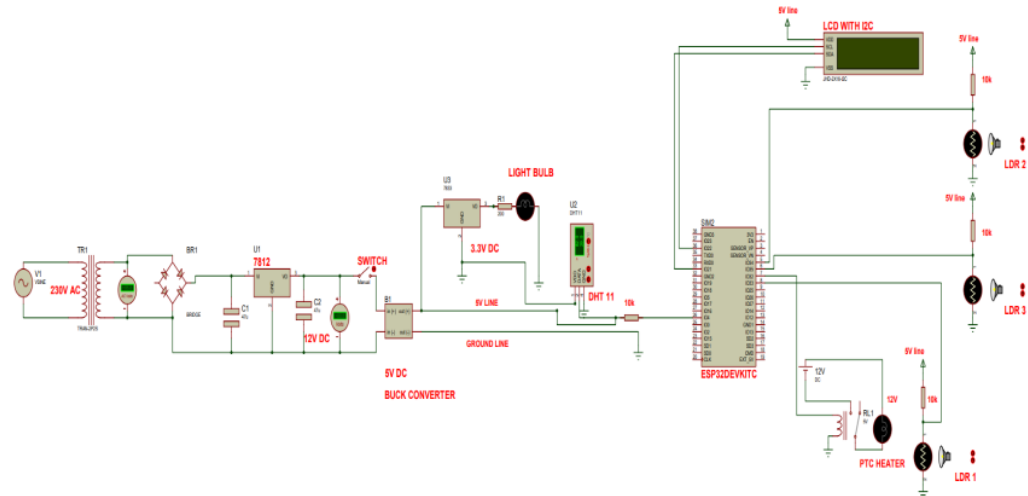


Figure 3.15: Circuit Diagram

3.3.3 Systems Flowchart

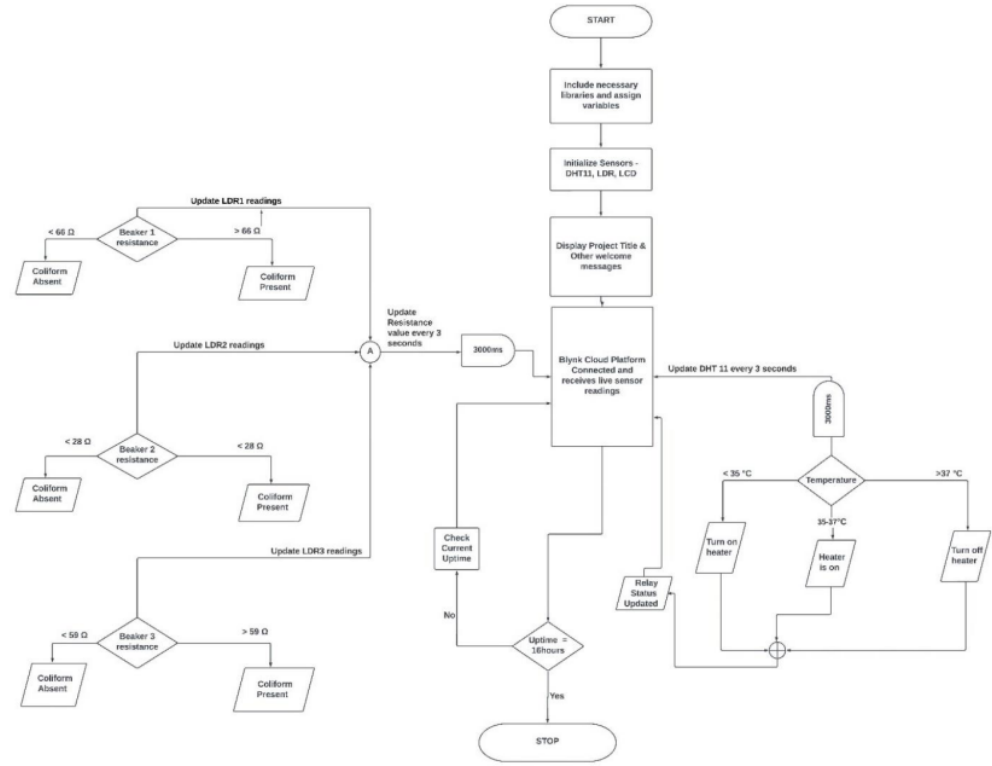


Figure 3.16: Systems Flowchart

3.3.4 Electrical Calculations

Heat transfer calculations

In order to determine the suitable power rating for the heating element necessary in the incubator, it is crucial to take into consideration the amount of heat energy required to elevate the temperature of both the water and air within the incubator to the desired level. By utilizing the specific heat capacity formula,

$$Q = m \times c \times \Delta\theta$$

8
3.1

Where:

Q is the amount of energy (in joules)

m is the mass of water (in kilograms)

c is the specific heat capacity of water (in J/kg.k)

$\Delta\theta$ is the change in temperature (in kelvin, K)

There are three beakers in total, each containing 100ml of water, making a total of

300ml. Converting 300ml to mass gives:

22
$$\text{Density} = \frac{\text{Mass}}{\text{Volume}}$$

$$\text{Mass} = \text{Density} \times \text{Volume}$$

Density of water = 1g/cm³ or 1g/ml, therefore, mass of water in incubator =

$$1\text{g/ml} \times 300\text{ml} = 300\text{g or } 0.3\text{kg}$$

4
The specific heat capacity of water is approximately 4190J/kg.k Assuming the initial temperature of the water is room temperature 20°C (293k), and we want to raise it to 37°C (310k), ΔT would be 17kelvin. Therefore, the heat energy required to raise the temperature of 100ml of water is:

$$Q = 0.1\text{ kg} * 4190\text{J/kg.k} * 17\text{k} = 7.123\text{kJ}$$

4
Since there are three beakers, the total heat energy required to raise the temperature of all the water is:

$$Q_{\text{total}} = 3 \times 7.123\text{kJ} = 21.369\text{kJ}$$

In addition to the water, it is important to factor in the heat energy needed to increase 13
the temperature of the air inside the incubator. To determine the quantity of heat

energy required for raising the temperature of a given volume of air, the following equation can be employed:

$$Q = m \times c_p \times \Delta\theta$$

79
Where:

m = mass of air

c_p = Specific heat capacity of air

$\Delta\theta$ = change in temperature

The incubator has internal dimensions of 270mm x 140mm x 200mm. Therefore, volume of incubator = Length x Breadth x Height = 0.27m x 0.14m x 0.2m = **0.00756m³**. We need to raise the temperature inside the incubator from 20°C (293k) to 35°C (308k). $\Delta\theta$ will be (**308k – 293k = 15kelvin**). The mass of air can be calculated using the density of air (**approximately 1.28 kg/m³**) and the volume of the incubator:

22
Density = Mass/Volume

Mass = Density x Volume

$$m = 1.28 \text{ kg/m}^3 * 0.00756\text{m}^3 = \mathbf{0.009677\text{kg}}$$

A
The specific heat capacity of air is approximately **1005 J/kg·K**. Therefore, the heat energy required to raise the temperature of the air inside the incubator is:

$$Q = 0.009677\text{kg} * 1005 \text{ J/kg·K} * 15 = \mathbf{145.88\text{J}}$$

34
The following amount of heat energy is needed overall to raise the temperature of the water and air inside the incubator:

$$Q_{\text{total}} = 21.369 \text{ kJ} + 145.88 \text{ J} = 21.514 \text{ kJ}$$

To calculate the power rating required for the PTC heating element, we also need to consider the time it takes to raise the temperature. Let's assume we want to raise the temperature in 20 minutes or 1200 seconds. Therefore, the power rating required for the PTC heating element is gotten by using the formula $Q = pt$

$$P = Q_{\text{total}} / t = 21.514 \text{ kJ} / 1200 = 17.9 \text{ watts appx } 18 \text{ watts}$$

Hence, a PTC element rated for about **18watts** will be sufficient to raise the ambient temperature in the incubator to 35-37°C in 20 minutes. A lower rating heating element could alternatively be used, as the polystyrene foam used to pad the walls of the incubator will enhance insulation.

COMPONENT	VOLTAGE RATING	CURRENT DRAW
ESP 32	5v	240mA
Relay	5v	160mA
Liquid Crystal Display (LCD)	4.5v - 5.5v	120mA
Heating Element	12v	1670mA
LED Bulb	3.4v	350mA
DHT 11	3.3v – 5.5v	2.5mA

Table 1: Component Voltage and Current Values

3.4 Description of Software Requirements

The programming of the entire system will be done using the Arduino C/C++ language on the Arduino Integrated Development Environment (IDE), followed by compilation and uploading to the microcontroller. The software required for this project is outlined below.

59

3.4.1 Arduino Integrated Development Environment (IDE)

This software is open-source and streamlines the process of programming and uploading code to the microcontroller board. It can be used on different operating systems such as Windows, Mac OS X, and Linux. The user interface was developed using Java and free and open-source tools like Lua Processing. Our project's Arduino board programming is written in C++, and the IDE software translates our readable human program codes into machine language. Before uploading to the microcontroller board, the IDE checks the source code for errors to ensure proper functionality.



60

Figure 3.17: Arduino Integrated Development Environment (IDE)

The screenshot shows the Arduino IDE interface with the following details:

- Title Bar:** Myron | Arduino IDE 2.3.0
- Tool Bar:** File Edit Sketch Tools Help
- Sketch Name:** Blynk
- Sketch Description:** ESP32 Dev Module
- Code Area:**

```

1  /*FINAL YEAR PROJECT -
2   TTIE: DEVELOPMENT OF A PORTABLE INCUBATOR FOR DETECTION OF COTIFORM USING IOT
3   RV: OLONISAKIN DAVID AKOLADE 18CKB24244 AND OGUUBE FAVOUR OZOMA 18CKB24232
4   EEE SET2023 #WITH SET
5   SUPERVISOR: DR. OHORUYI OSEHWAINGIE */
6
7 //Global declarations//
8 //Including necessary libraries
9 #include <WiFi.h>
10 #include <ESPClient.h>
11 #include <BlynkSimpleEsp32.h>
12 #include <LiquidCrystal_I2C.h>
13 #include <Wire.h>
14 #include <DHT.h>
15
16 //Blynk and wifi details
17 #define BLYNK_TEMPLATE_ID "TMPLID4_oZig"
18 #define BLYNK_TEMPLATE_NAME "COTIFORM DETECTION"
19 #define BLYNK_AUTH_TOKEN "O_7ewrj-zfD9Y49K4k0Tx3uwyJut6y"
20 #define BLYNK_PRINT Serial
21 //wifi credentials and assignments
22 char auth[] = BLYNK_AUTH_TOKEN;
23 char ssid[] = "Cotiform";
24 char pass[] = "bacteria";
25 const long startTIme;
26 BlynkTimer timer;
27 LiquidCrystal_I2C lcd(0x27, 20, 4); //20x4 lcd dinmsplay
28 // initialize DHT sensor
29 #define DHTPIN 5
30 #define DHTTYPE DHT11

```
- Output Area:** Serial Monitor K
 - Not connected. Select a board and a port to connect automatically.
- Status Bar:** In 47, Col 48 - ESP32 Dev Module (not connected)

Figure 3.18: Arduino Integrated Development Environment (IDE) Code Snippet

3.4.2 Blynk

Blynk is a versatile platform that enables users to create mobile applications that control and monitor devices over the internet. One of the key advantages of Blynk is its user-friendly interface, which makes it easy for users to monitor and control various IoT devices from their smartphones. In this particular project, Blynk offers a significant role in facilitating remote monitoring of the incubator's temperature and water quality parameters using a mobile device.

By integrating the ESP 32 with the Blynk platform, users can remotely monitor the temperature of the incubator and the water quality parameters in real-time. Blynk also allows users to set up alerts and notifications that can be sent to their smartphones if the temperature or water quality parameters fall outside the acceptable range. This feature ensures that users can respond quickly to any deviations from ideal conditions and prevent damage to the organisms being cultured in the incubator.

In summary, Blynk's user-friendly and convenient interface makes it an ideal platform for remotely monitoring and controlling the incubator's temperature and water quality parameters. By using Blynk in conjunction with the ESP 32, users can ensure that optimal conditions are maintained for the organisms being cultured in the incubator.

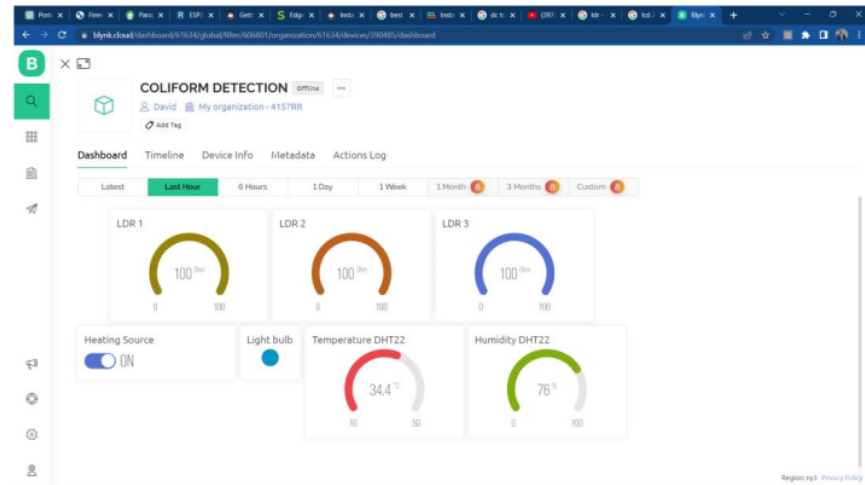


Figure 3.19: Blynk Application

3.5 Conclusion

This section provides a detailed overview of the construction process of this project. It explains in detail the different components utilized, the circuitry, block diagrams, algorithms, and flowcharts employed. Moreover, it describes the architecture and techniques of the system, giving practical examples. The chapter also outlines the implementation procedures, including the proposed model and the development approach to be followed.

CHAPTER FOUR

RESULTS AND DISCUSSIONS

4.1 Introduction

This chapter presents the outcomes and analysis of the project, focusing on significant discoveries and observations made during the implementation and testing phases. The primary objective of the project was to develop an incubator system that could detect coliform in water by controlling temperature and humidity within a predetermined incubation period. To accomplish this, the incubator utilized several components, such as an LDR sensor, a DHT sensor, a PTC heating element, an ESP 32, and the Blynk IoT platform.

The chapter details the interconnection of the hardware components and provides a step-by-step account of calibrating the LDR sensor. Accurate readings from these sensors are crucial for drawing meaningful conclusions. It also covers the configuration of the PTC heating element and the integration of the hardware components with the Blynk platform.

Moreover, comprehensive testing of the individual subunits forming the complete system was conducted to identify any design flaws that might have arisen during the project's construction. The testing phase aimed to ensure the proper functioning of the system while rectifying any potential errors.

4.2 Hardware Implementation

The interior structure of the incubator and the connectivity of all project-related components are both crucial aspects. To ensure optimal thermal insulation, the inside walls of the incubator were lined with polyethylene foam. This choice was made because polyethylene foam possesses excellent properties for thermal insulation. Its low thermal conductivity effectively inhibits heat transfer, minimizing heat loss to the surroundings. By implementing this insulation, the energy needed to maintain the desired temperature range is reduced, allowing for a consistent and regulated interior temperature.

CHAPTER FIVE

CONCLUSION

5.1

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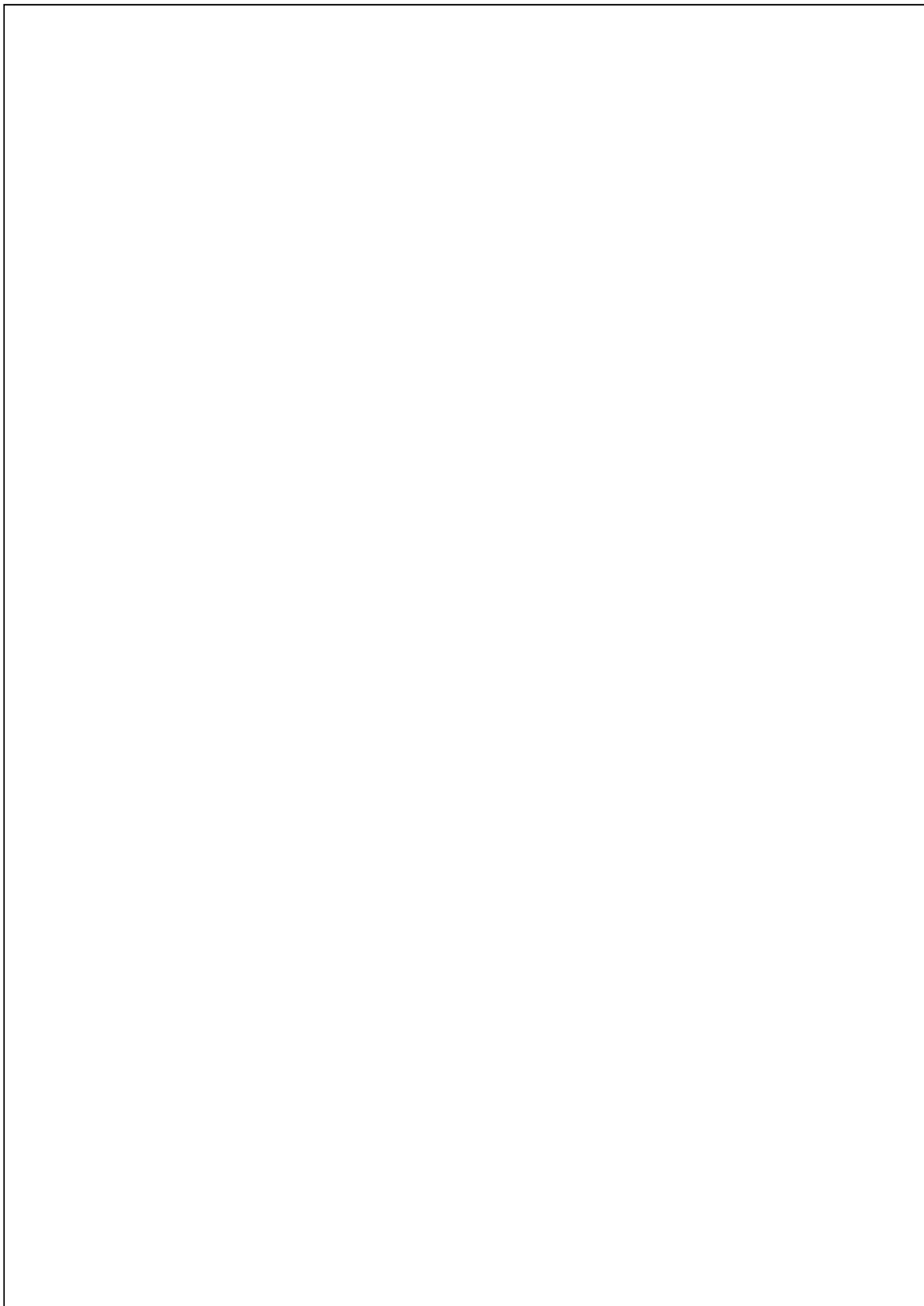
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