

Original Research Article

Protective Effect of Methanolic Extract of *Vernonia amygdalina* (Bitter-Leaf) on Alloxan-Induced Pancreatic Toxicity In Adult Wistar Rats

ABSTRACT

The pancreas is a glandular organ endowed with two main functions; an exocrine function that assists digestion and an endocrine function that modulates blood glucose concentration. Pancreatic Beta-cells produce insulin to regulate/lower blood glucose concentration, driving them into cells to be utilized. Distortions of the architecture of pancreatic Beta-cells lead to function loss; thus, preservation/restoration of their cytoarchitecture would sustain their functions. *Vernonia amygdalina* is a vital plant acknowledged widely for its antioxidant and anti-diabetic effects; however, its impact on the pancreas has received little attention. This study is premeditated to examine the effects of the methanolic leaf extracts of *Vernonia amygdalina* (MLEVA) on the cytoarchitecture of the pancreas in Alloxan-induced pancreatic toxicity. Thirty (30) adult male Wistar rats were grouped into 6 (A-F) (n=5). Group A, serving as the control group, received feed and water only. Group B received a single intraperitoneal injection of 150mg/kg of Alloxan. Groups C, D, and E received single intra-peritoneal injections of 150mg/kg of Alloxan and were then treated with 200, 300, and 400 mg/kg/day of the MLEVA orally, respectively. Group F received a single intraperitoneal injection of 150mg/kg of Alloxan and was treated with 100mg/kg/day of Vitamin E orally. The experiment lasted for 16 days before sacrificing the animals via a median incision on the abdominal cavity, under ketamine (100mg/ml), as an anesthesia, 24 hours after their last treatment. The pancreas was carefully and rapidly processed for routine H & E staining. This study recorded a 28% mortality rate after Alloxan administration and led, leading to degenerative histo-pathological changes in the pancreas' endocrine and exocrine compartments. 300mg/kg of MLEVA demonstrated a marked regenerative effect on the pancreas. The MLEVA is a promising agent for the management

~~of managing diabetes~~Diabetes, evidenced by its protective and therapeutic effect on Alloxan-induced pancreatic toxicity.

Keywords: *Vernonia amygdalina*, MLEVA, Pancreatic toxicity, Alloxan, Diabetes Mellitus

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INTRODUCTION

The ~~Panereas~~pancreas is a glandular organ located retroperitoneally within the abdominal cavity.^[1] It is a gland that is partly exocrine and partly endocrine. The exocrine part secretes enzymes that assist in the digestion of carbohydrates, proteins and fats, ~~while~~. At the same time, the endocrine pancreas produces two major hormones in addition to Somatostatin directly into the portal circulation.^{[1][2]} These hormones include; glucagon secreted by the alpha-cells and insulin secreted by the beta-cells. The exocrine pancreas has a compound tubuloalveolar appearance microscopically. However, the endocrine pancreas appears as numerous rounded collections of cells embedded within the endocrine part. They are known as islets of Langerhans or ~~simply~~termed pancreatic islets.^[1] ~~Therefore~~ Therefore the function of the pancreas can easily be evaluated by examining the activities of these pancreatic hormones. The major hormones produced by the endocrine pancreas functions by regulating glucose.^[2] Glucose is the main sugar found in blood, serving as the ~~body's~~ body's source of immediate energyimmediate energy source. Notwithstanding, ~~regulating~~regulation of blood glucose levels is vital for the healthy living of any complex organism. The ~~Panereas~~pancreas is a vital organ charged mainly with the responsibility of blood sugar regulation, amongst other functions. Glucagon promotes an increase in ~~the~~blood glucose concentration, while insulin counters this action, by enhancing the uptake of glucose by cells ~~in order~~to be ~~utilised~~utilized.^{[2][3]} The pancreas ~~however, just like other organ, is prone to toxicities or degenerative diseases which, however, is prone to toxicities or degenerative diseases that~~ can be life-threatening. Ailments such as pancreatic acinar atrophy, pancreatitis and Pancreatic autoimmune disease (which leads to Insulin-dependent diabetes mellitus) diminishes the functionality of the pancreas, and, if left untreated, can prove lethal.^{[2][3]} The life-

threatening effect of pancreatic pathologies is mostly ~~characterised~~ characterized by the loss of blood glucose concentration regulation, which can easily tip an individual into ~~Diabetis-Diabetes~~ Mellitus and its complications.

In Africa, ubiquitous herbal medications have been attributed curative elements to ~~Diabetes-Diabetes~~ Mellitus.

Vernonia amygdalina, a plant commonly seen across the west coast of Africa where it is cultivated and ~~also~~ grows wild as a domestic browse plant, has been attributed such a curative element.^[4] ~~It is commonly identified as "bitter leaf" in Nigeria~~ Nigeria commonly identifies it as a "bitter leaf" because the leaves and the stem have an astringent, bitter taste. In Nigeria, it is a major vegetable of the celebrated ~~"bitter leaf soup"~~ "bitter leaf soup". *Vernonia amygdalina* has a long history of use in folk medicine, particularly among the people of sub-Saharan Africa and Southern Nigeria, ~~where it~~ It has a high reputation for use in the traditional management of diabetes mellitus.^{[4][5][6]} Scientific studies have also reported its antihyperglycemic action^{[7][8]} and hypoglycemic action^[9] in diabetic and non-diabetic rats, respectively. The methanolic leaf extract has been shown to possess anti-diabetic and antioxidant Potentials in Alloxan-Induced ~~diabetis~~ diabetes mellitus in Sprague-Dawley Rats.^[10] However, despite the extensive usage of *Vernonia amygdalina* traditionally as well as experimentally as a remedy for diabetes mellitus, its therapeutic potential is still largely unexplored, predominantly when it comes to its effect on organ and tissue architecture. Therefore, this study aims to investigate the effects of the methanolic extract of bitter-leaf (*Vernonia amygdalina*) on the microstructure of the pancreas in Alloxan-induced pancreatic toxicity using rat models.

Alloxan is a toxic glucose analogue that preferentially accumulates in the pancreatic beta cells via the GLUT2 glucose transporter.^{[11][12]} This initiates a series of cyclic redox reactions, generating reactive oxygen species and hydroxyl radicals that lead to the death of beta cells.^[12] It is commonly used in the induction of experimental ~~diabetes-Diabetes~~ giving rise to the rat model of insulin-insulin-dependent diabetes mellitus.^[13] It also causes an enormous reduction in glucose-induced insulin release by inhibiting the β -cells of the islets of Langerhans glucose sensor glucokinase, thus inducing hyperglycemia.^{[14][15][16]} Alloxan dosages ranging from 35mg/kg to 90mg/kg

have been used experimentally to induce a rat model of ~~insulin-insulin~~-dependent diabetes mellitus.^{[11][17]} However, increased doses of Alloxan leads marked beta-cell toxicities and remarkable damage in the endocrine pancreas.^{[18][19]} It also damages the exocrine pancreas.^{[20][21]}

Although endocrine evaluation serves as a direct approach ~~in to~~the monitoring of pancreatic toxicity, the use of histology in rat models serves as a holistic tool.^[22] Distortion of the cyto-architecture of pancreatic acini, duct system and surrounding endocrine cells is a marker of pancreatic toxicity, hence used in the demonstration of the study of *Vernonia amygdalina* effect on ~~distorted-distorted~~ pancreatic cyto-architecture.

MATERIALS AND METHODS

Plant Materials

Matured bitter leaves (*Vernonia amygdalina*) were acquired from a local farm in Enugu-South local government area, Enugu state. ~~These leaves were identified and authenticated by a taxonomist at the Faculty of Agricultural Science, Enugu State University of Science and Technology~~A taxonomist at the Faculty of Agricultural Science, Enugu State University of Science and Technology, identified and authenticated these leaves.

Processing of Plant Materials

The fresh bitter leaves were washed and air-~~dry-dried~~at room temperature for one week within the animal facility where the study took place. The dried leaves were macerated using a warren blender to a smooth, dry powder and ~~was-were~~subjected to methanolic extraction. The methanolic extraction technique used was adopted from Oyedeji et al., (2013)^[23] and Adefisayo et al., (2018)^[24]. filtered with Whatman No.1 filter paper (150 mm). The extract was concentrated using a rotary evaporator at 40–50 °C under reduced pressure. 10g of the crude extract was diluted in 100ml of normal saline and stored in an airtight container afterwards. The extract was stored at –8°C until required for use.

Experimental animals

Thirty adult male Wistar rats with average weights of 180-200g were procured from the animal house facility of the Department of Anatomy, University of Nigeria, Enugu campus. However, this study was carried out in the ~~Animal~~ facility of the Enugu State University of Science and Technology College of Medicine, Parklane, Enugu. The animals were kept in ~~well~~-ventilated breeding rooms and ~~housed~~ in netted iron cages. ~~There were provided easy access to water and standard livestock pellets (Guinea Feed Nigeria Limited) as food, and were also~~ Easy access to water and standard livestock pellets (Guinea Feed Nigeria Limited) was provided as food, and they were allowed to acclimatize for 2 weeks. The animals were maintained under standard laboratory conditions, and handling was done following international guidelines on the use of experimental animals.

Experimental design

The rats were randomly divided into six (6) groups (A-F) of 5 rats each. Alloxan was administered ~~via intraperitoneal route~~ intraperitoneal, while all extract administrations were done orally. The experiment lasted 16 days.

Group A served as the control group and received normal saline orally, till the end of the experiment. Group B ~~served~~ was the untreated positive control group and received only a single intraperitoneal administration of 150mg/kg ~~bwt~~ bwt of Alloxan. This dosage of Alloxan used was adopted from Cheekati et al., (2017).^[19] Groups C, D and E received a single intraperitoneal administration of 150mg/kg of Alloxan, left for two days (48 hours) to confirm diabetic statuses and then treated with 200, 300 and 400 mg/kg/day of the methanolic extract of bitter-leaf (*Vernonia amygdalina*) respectively for 14 days. The extract dosage was adopted from Adefisayo et al., (2018).^[24] Group F received a single intraperitoneal administration of 150mg/kg of Alloxan, left for two days (48 hours) to confirm the diabetic status and then treated with 100mg/kg/day of Vitamin E for 14 days.

Confirmation of diabetic status:

All animals were deprived of both food and water for 12 hours. Under aseptic conditions, blood samples were collected on the first day from all groups via tail ~~venopuncture~~ venipuncture, and their blood glucose was determined using a ~~one-one-~~ touch glucometer strip test. Blood glucose levels for all groups ranged from 97mg/dl to 115mg/dl. Group B-F was administered with a single intraperitoneal administration of 150mg/kg of Alloxan immediately after ~~the determination of~~ determining their blood glucose. ~~To avoid mortalities due to hypoglycemic shock after the induction, their drinking water was replaced with an oral solution of 20% glucose for two days~~ After the induction, their drinking water was replaced with an oral solution of 20% glucose for two days to avoid mortalities due to hypoglycemic shock.^[25]

Blood samples were re-collected from group B-F via tail venipuncture after two days (48 hours), and their blood glucose was determined. Blood glucose levels for these groups after 48 hours ranged from 418mg/dl to 505mg/dl, confirming positive hyperglycemia and suggesting pancreatic toxicity. Previous studies by Akhtar et al., (2002)^[26] and Díez et al., (2013)^[27] have reported that animals exhibiting a significant elevation in blood glucose levels above 250 mg/dL should be considered diabetic.

Histological Study

The animals were sacrificed 24 hours after the last administration via a median incision on the abdominal wall, under ketamine (100mg/ml), as anaesthesia. The pancreas was hurriedly isolated by dissection and immediately fixed with 10% formalin in labelled containers for 72 hours ~~prior to~~ before processing. The fixed tissues were processed using the standard protocols for histological tissue processing and stained with hematoxylin and eosin for histological studies. Photomicrographs were taken using Amscope 14MP USB 3.0 digital microscope camera at x200 magnification.

RESULTS

Animal mortality

2 out of 5 animals died from Group B ~~which served as~~, the untreated positive control group. 1 out of 5 animals each died from Group C and E being treated with 200 mg/kg

and 400 mg/kg of the methanolic extract of *Vernonia amygdalina*, respectively. 3 out of 5 animals died from Group F being treated with 100mg/kg of Vitamin E. Therefore, 7 out of 25 rats administered with Alloxan died before the ~~end of the experiment~~ giving rise to experiment's end, giving rise to a 28% mortality rate.

UNDER PEER REVIEW

Histological Analysis

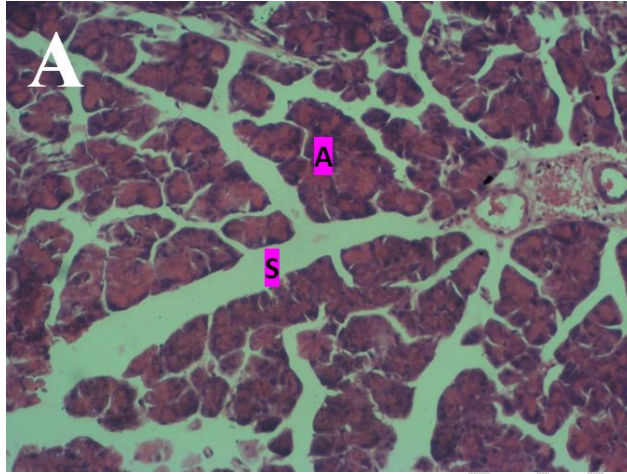


Fig 1.(Photomicrograph of a section of the pancreas of the control animal group fed with only food and water showing the normal acini tissue (a) with septa (s).General tissue appears normal. H&E.x200)Group A

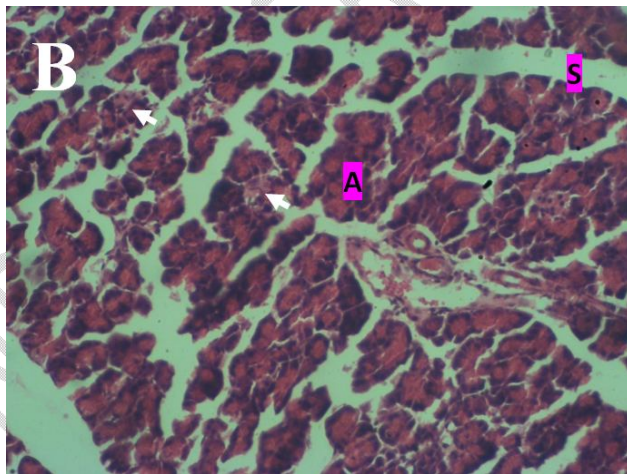


Fig .2 (Photomicrograph of the pancreas of the **untreated** animal group administered only a single intraperitoneal administration of 150mg/kg/bwt of Alloxan, showing acini metaplasia, dilated blood vessels and marked shrinkage of the pancreatic islets (**arrow**) with leukocyte infiltration. H&E.x200)Group B

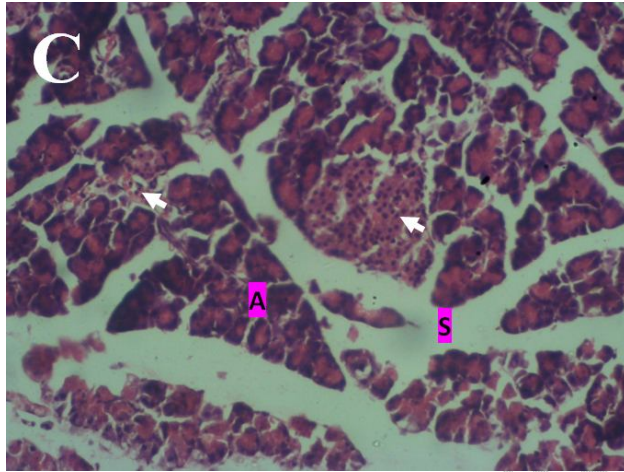


Fig .3: (Photomicrograph of the pancreas of the animal group treated with 200mg/kg of the methanolic extract of *Vernonia amygdalina* showing focal areas of mild acini aplasia and mild leukocyte infiltration in the pancreatic islets (arrow). Acini tissue (a), septa (s). H&E.x200) Group C

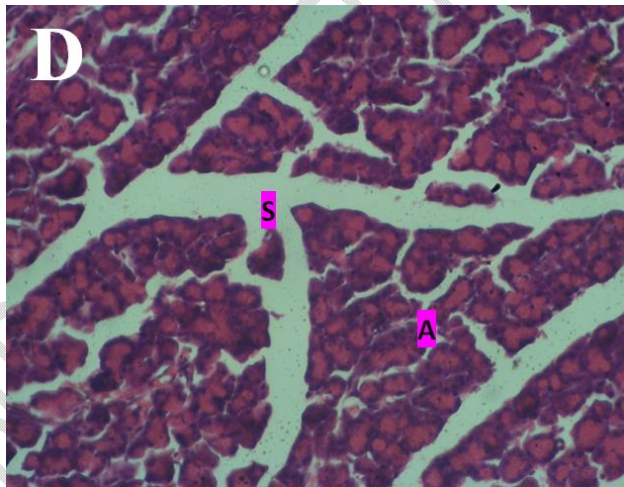


Fig .4: (Photomicrograph of the pancreas of the animal group treated with 300mg/kg of the methanolic extract of *Vernonia amygdalina* showing normal acini tissue (a) with septa (s). General tissue appears normal. H&E.x200) Group D

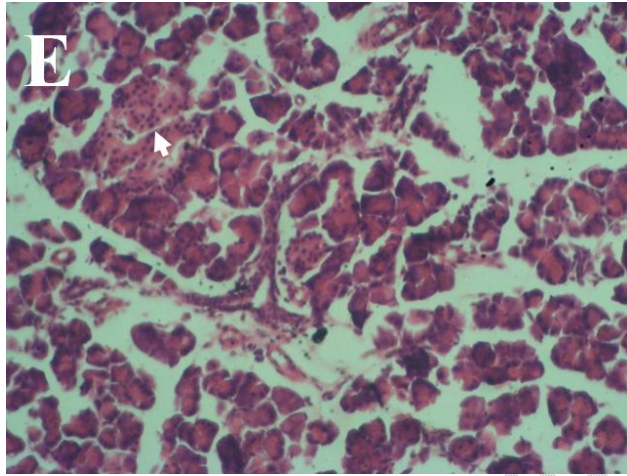


Fig .5:(Photomicrograph of the pancreas of the animal group treated with 400mg/kg of the methanolic extract of *Vernonia amygdalina* showing mild tissue derangement with necrotic and degenerative changes in the pancreatic islets. H&E.x200)Group E

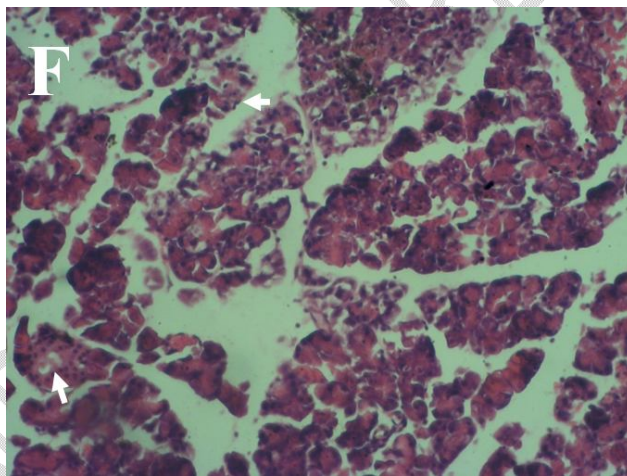


Fig.6: (Photomicrograph of the pancreas of the animal group treated with 100mg/kg of vitamin E showing mild acini derangement with fibrosis and atrophy of pancreatic islets. H&E.x200)Group F

DISCUSSION

Animal mortality has previously been reported as a complication of Alloxan injections. Previous studies have indicated that Alloxan administration leads to characteristic blood glucose responses, including a severe, transitional hypoglycemia that may be fatal.^[12] However, the data from previous studies are inconsistent, as some researchers reported mortality rates of up to 75% with an Alloxan dose of 80 mg/kg,^[28] ~~In comparison, while~~ others reported 100% survival rates with a ~~dose of 150 mg/kg~~ 150 mg/kg dose.^[29] ~~In the midst of~~ Amid these inconsistent results, reported death rates differ substantially depending on the animal species, dose, chemicals used, and protocols employed.^[25]

This study recorded a 28% mortality rate after Alloxan administration. This low value is suggested to be due to ~~the replacement of~~ replacing their drinking water with an oral solution of 20% glucose for two days. Bacevic et al., (2020)^[25] indicated that this practice helps avoid mortalities due to hypoglycemic shock after Alloxan induction.

A single intraperitoneal administration of 150mg/kg/bwt of Alloxan led to degenerative histopathological changes ~~on in both the endocrine and exocrine compartments of the pancreas~~ the pancreas endocrine and exocrine compartments characterized by acini metaplasia and marked shrinkage of the pancreatic islets with leukocyte infiltration. This is ~~in accordance with~~ per previous studies by Cheekati et al., (2017),^[19] who reported that a single dose of 150 mg/kg of Alloxan intraperitoneally resulted ~~to in~~ degenerative changes, followed by fibrosis and atrophy of islets and other pancreatic tissues. El-Esawy et al., (2016)^[30] also demonstrated that intra-peritoneal injections of 150 mg/kg Alloxan led to both endocrine and exocrine pancreatic tissue damage characterized by degeneration and vacuolizations in islet cells, congestion and dilation of blood sinusoid, widening of the intercalated duct and the formation of fibres periphery to dilated blood vessels and ducts. The degenerative changes of Alloxan administration ~~is are~~ attributed to the inhibition of insulin secretion from the pancreatic beta cells via beta cell toxicity^[12] and possibly through the mechanism of induction of free radical species^[13] and oxidative stress within the pancreatic tissue.

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Previous studies have reported that ~~intake of vitamin E prevented Alloxan induced pancreatic necrosis by preventing severe oxidative stress, strongly enhancing insulin secretion from the pancreas and improves the incidence of hyperglycemia~~ intake prevented Alloxan-induced pancreatic necrosis by preventing severe oxidative stress, strongly enhancing insulin secretion from the pancreas and improving hyperglycemia incidence.^[31] However, this study revealed that treatment with 100mg/kg of vitamin E still demonstrated mild degenerative changes ~~both~~ in the endocrine and exocrine compartments of the pancreas characterized by mild acini derangement with fibrosis and atrophy of pancreatic islets. This finding suggests that Vitamin E might ~~prove to~~ be more therapeutic as a preventive remedy as it displayed the therapeutic insufficiency of 100mg/kg of vitamin E in curing/regenerating the pancreatic cells after ~~Alloxan~~ Alloxan-induced toxicity. Nevertheless, Malek et al., (2010)^[32] classified vitamin E as a strong antioxidant, but its efficacy depends on the used dosage of vitamin E and the time length of administration.

Treatment with the ~~methanolic leaf extracts of Vernonia amygdalina (MLEVA) showed dose depend~~ Vernonia amygdalina (MLEVA) 's methanolic leaf extracts showed dose-dependent therapeutic potentials. The findings from this study suggest ~~low~~ low-dose insufficiency and ~~high~~ high-dose toxicity. 200mg/kg and 400mg/kg of MLEVA still demonstrated mild degenerative changes ~~both~~ in the endocrine and exocrine compartments of the pancreas characterized by mild acini aplasia, mild tissue derangement and also necrotic pancreatic islets. However, 300mg/kg of MLEVA demonstrated a marked regenerative effect as the pancreatic tissues displayed normal cyto-architecture on both the endocrine and exocrine compartments of the pancreas. This study attributes this potential to the phytochemical constituents of the plant extract. Adefisayo et al., (2018),^[24] reported a high amount of flavonoids and phenolic contents found in the methanolic leaf extract of *Vernonia amygdalina* after the quantitative phytochemical screening. These contents are responsible for its antidiabetic^{[7][33][34][35][36]} and antioxidant potential^{[37][38][39]} formerly described by previous studies.

CONCLUSION

This study investigated the effects of the methanolic extract of bitter-leaf (*Vernonia amygdalina*) on the microstructure of the pancreas in Alloxan-induced pancreatic toxicity using rat models. Low doses displayed insufficiency in producing the expected results, while high doses worsened Alloxan-induced toxicity. However, 300mg/kg of MLEVA demonstrated a marked regenerative effect on the cyto-architecture of ~~both the endocrine and exocrine compartments of the pancreas~~ pancreas's endocrine and exocrine compartments. The MLEVA is a promising agent for the management of Alloxan-induced pancreatic toxicity. Nevertheless, caution should be applied while consuming this plant.

CONSENT

It is not applicable

ETHICAL APPROVAL

Ethical clearance was obtained from the Research and Ethical Clearance Committee, Faculty of Basic Medical Sciences, College of Medicine, Enugu State University of Science and Technology with ethical clearance code **ESUCOM/FBMS/ETR/2022/027**

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