

Bharath

INSTITUTE OF HIGHER EDUCATION AND RESEARCH

(Declared as deemed to be university under section 3 of UGC Act 1956, vide notification No.F.9-5/2000-U.3)

25.10.19

From
Head of the Department,
Department of Genetic Engineering,
Bharath Institute of Higher Education and Research,
Chennai

To
The Dean Engineering,
Bharath Institute of Higher Education and Research,
Chennai

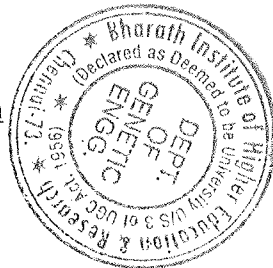
Respected Sir,

Sub: Requesting for Industrial visit- Reg

I would like to request you to permit our 2018 Batch Genetic Engineering students for Industrial visit. I have enclosed the Student name list for your reference. We request you to approve the Industrial visit in the ARMATS Biotech Labs Pvt.Ltd, Chennai and do the necessary transportation facility for the benefit of students and staffs.

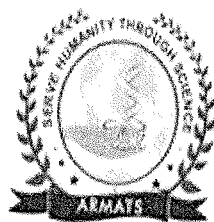
Thanking you

Dr. P.B Ramesh babu
Email- rameshbabu.bi@bharathuniv.ac.in
Phone- 9841788001



Yours sincerely

Dr. P.B. RAMESH BABU, Ph.D.,
Professor and Head
Department of Genetic Engineering,
Education - Research
(Declared as deemed to be university under section 3 of UGC Act 1956)



ARMATS BIOTEK
TRAINING AND RESEARCH INSTITUTE (ABTRI)

To

The Head of the Department,
Department of Genetic engineering,
BIHER.

Sir,

Sub: Req for Industrial visit – Reg

Referred to letter dated 25.10.2019 regarding Industrial visit we permit your students for Industrial visit at Maduvankarai, Guindy, Chennai 600032 on 30.10.2019

Regards,

Dr. Arumugam

Register office:

New 14/18, Mettu street/Link road

Maduvankarai, Guindy

Chennai 600032

Students Industrial Visit Report - Armats Biotek Training and Research Institute

The B.Tech II year students of Genetic engineering, BIHER had visited the Institute and known about following processes:

- Electrophoresis – SDS PAGE
- Polymerase Chain Reaction
- ELISA – Direct ELISA

Detailed report of the Industrial visit

Polymerase Chain Reaction:

PCR theoretically amplifies DNA exponentially, doubling the number of target molecules with each amplification cycle. The objectives of this experiment involved to Isolate the DNA genome ,amplify a specific gene and to understand the principles of the polymerase chain reaction. The DNA strands were unwound (denatured) by enzyme to form two single strands. A RNA polymerase binds and synthesizes a short complementary piece of RNA on the DNA strand at the initiation site of replication. Followed by the DNA hetero duplex acts as a priming site for the DNA polymerase that binds and produces the complementary strand. Totally three steps of the PCR was performed which includes majorly, First, the template DNA or genetic material was denatured; the strands of its helix were unwound and separated-by heating to 90-96°C. The second step is hybridization or annealing. The Taq polymerase was added to initiate DNA replication, with a few length RNA primers. In the PCR reaction, short complimentary double stranded oligos primers which were complimentary to the gene sequence were added that bind the denatured DNA and act as origins of replications. Following denaturation, the reaction mixture was rapidly cooled to a temperature below the melting point of the specific primers (~55°C). In the third step, the temperature of the reaction was gradually raised to the optimal temperature for the polymerase (68-72°C). The polymerase synthesized new DNA, starting from the primer, the polymerase reads a template strand and generated complementary nucleotides very quickly. The result was two new helices in place of the first, each composed of one of the original strands plus its newly assembled complementary strand.

Electrophoresis – SDS PAGE

The different buffers required for SDS-PAGE were prepared and the Casting gels for SDS-PAGE were prepared. The surface of the balance with a tissue paper was cleaned and a butter paper was placed on the balance and tare the weight of paper before weighing. 0.5 M Tris-HCl pH 6.8 and 1.5 M Tris-HCl pH 8.8 were prepared according to the protocol. tris-base was dissolved in minimum volume of water and proceeded for adjusting the pH of the buffer to the required value. The pH of 0.5 M tris base solution was adjusted to 6.8 using concentrated HCl. The pH of 1.5 M tris base solution was gradually adjusted to 8.8 using concentrated HCl. 144.1g of glycine, 30.3g of tris base and 10g SDS was weighed and dissolved in distilled water. For the Buffer preparation, the components on a magnetic stirrer were dissolved. For the Gel casting 10% SDS and 10% APS was prepared by weighing the required amount and dissolving it in water. 12.5% resolving gel solution was prepared, consisting of 3ml acrylamide and bis-acrylamide, 1.5ml of 1.5M Tris-HCl pH 8.8, 60µl of 10% SDS and 60ul of 10% APS, 2.11µl TEMED and 1.38ml water. 12.5% resolving gel prepared earlier slowly was poured along the sides of the glass plates. Stacking gel was prepared consisting of 1 ml acrylamide and bis-acrylamide 1.8ml 1.5M Tris-HCl pH- 6.8, 75ul of 10% SDS and 75ul of 10% APS, 4.5ul TEMED and 1.6ml water. A comb was inserted inside the stacking gel immediately after its transfer into the assembly and the gel was allowed to polymerize for 25 minutes. The combs were removed and the samples were loaded into the wells and buffer was loaded and the electrophoretic run was proceeded. The power-pack was switched OFF, the lid was removed and the glass plates were taken out of the electrophoresis unit. Finally the gel was carefully removed and place it in a tank containing water for further processing.

ELISA – Enzyme linked Immuno sorbent assay

Direct ELISA :

Antigen Coating was the first procedure in which the purified antigens solution was diluted to a final concentration of 1-10 µg/ml in bicarbonate/carbonate antigen-coating buffer (100 mM NaHCO₃ in deionized water; pH adjusted to 9.6). 100 µL of diluted antigen was added to each well of a microtiter plate. The plate was covered with adhesive plastic and incubated at 4°C overnight (or 37°C for 30 min) and the coating solution was removed and the plate was washed 3X with 200 µL PBS (Phosphate Buffered Saline) buffer (10 mM Na₂HPO₄ and 1.8 mM NaH₂PO₄ in deionized water with 0.2% Tween 20; pH Adjusted to 7.4) with for 5 minutes each time. The second procedure involved Blocking in which 200 µL blocking buffer (5% w/v non-fat dry milk in PBS buffer) was pipetted per well to block residual protein-binding sites. The plate was covered with adhesive plastic and incubated for 1-2 hour(s) at 37°C. The blocking solution was removed and and the plate

was washed 2X with 200 μ L of PBS for 5 min each time. The plate was flicked and pat dry. The third step was Reagent Preparation, the diluted standard solutions were prepared followed by the Primary Antibody Incubation. Serially diluted the conjugated primary antibody with blocking buffer immediately before use. 100 μ L of diluted secondary antibody solution was pipette into each well and the plates were covered with adhesive plastic and incubated for 2 hours at room temperature and finally the content in the wells was removed and washed them 3X with 200 μ L PBS buffer for 5 min each time. For the Substrate Preparation, the substrate solution was prepared immediately. The enzymes used for signal detection were horse radish peroxidase (HRP) and alkaline phosphatase (AP), and their corresponding substrates, stopping solutions, detection absorbance wavelengths and colour developed. For the Signal Detection step, 90 μ L of substrate solution was pipette in to the wells with the control and standard solutions and the Incubation of the plate at 37°C in the dark was done. Shades of blue was observed in the wells with the most concentrated solutions. Other wells showed no obvious color. 100 μ L of stopping solution to the wells was pipetted. The sample concentration from the standard curve was interpreted.

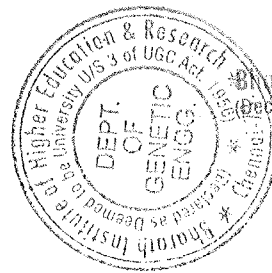
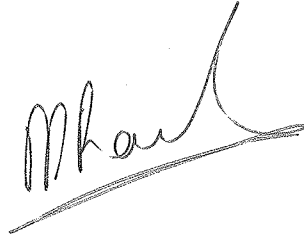


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STUDENTS NAME LIST

U19BR001	U19BR002
U19BR002	ALFRED D THEKKATH
U19BR003	AMBATI SANDEEP KUMAR REDDY
U19BR004	ARUMILLI LAKSHMI SARVANI
U19BR005	BODLA SAI HARSHA
U19BR006	CHANDRU J
U19BR007	DESAI JAYDEEP SARJERAO
U19BR008	FARHANA MUMTAJ A
U19BR009	GAYATHRI R
U19BR010	GOKARNAM ADARSH
U19BR011	GOKULAKRISHNAN V
U19BR012	GOVARDHAN R
U19BR013	HARI KRISHNAN V R
U19BR014	HARI PRIYA J S
U19BR015	HEMA DHARSHINI S
U19BR016	JAYA PRIYA V
U19BR017	KALLEPU SAI KUMAR
U19BR018	KAMALESH R
U19BR019	KEERTHI P
U19BR020	KURAPATI JAGADEESWAR REDDY
U19BR021	KUZHALI A
U19BR022	LANCY L
U19BR023	LATHIF A K
U19BR024	LAVANYA K
U19BR025	MAHARAJAN R
U19BR026	MARIYA BANU SRI A R
U19BR027	NAGALAKUNTA SINDHU
U19BR028	NAKULAN R K
U19BR029	NANDINI A I
U19BR030	NIKHIL ROY ANTHONY
U19BR031	NIMI SANTHU
U19BR032	NITISH KUMAR BHANDARI
U19BR033	NUNE SARATH KUMAR
U19BR034	PAPANI ANUSHU
U19BR035	POORANI V T
U19BR036	PRIYANKA S
U19BR037	PULLURU DIVYA DARSHINI
U19BR038	RAKSHANA G
U19BR039	RAMUGALLA PRANAY
U19BR040	REDDI SHIVA KUMAR
U19BR041	RITHUPARNA K

U19BR042	SACHIN SUBASHINI PM
U19BR043	U SAKTHI PRIYA
U19BR044	SHANIVARAPU VENKATARAJASEKHAR REDDY
U19BR045	SONALI KUMARI
U19BR046	SRINIVASAN V
U19BR047	SURYA G
U19BR048	SUSHMITHA K
U19BR049	TENUJ CHAUHAN
U19BR050	THOKCHOM CHRISTINA
U19BR051	K P TINAA SHREE
U19BR052	BARATHKUMAR B



Dr. P.B. RAMESH BABU, Ph.D.,
Professor and Head
Department of Genetic Engineering,
Shaoh Institute of Higher Education & Research
(Declared as Deemed to be University U/S 3 of UGC Act, 1956)
Chennai-600 073. INDIA