INTRODUCTION

Tuberculosis is an important infectious disease of human beings from the time immemorial. In the modern times it has been seen to re-emerge with drug resistance. Mycobacterium tuberculosis is responsible for infecting 10 million cases each year. Approximately 2 million people die of tuberculosis each year. In Pakistan; it is a major health problem. According to WHO report, it shows a total prevalence of 376/lack/year for the year 2012.

Mycobacterium tuberculosis commonly infects human lung but extra-pulmonary tuberculosis is not less common and infects almost all tissues of the body. Importantly tuberculosis in all locations is equally serious and even fatal.

Mycobacterium tuberculosis is a member of M. tuberculosis complex which comprises of 9 closely related strains that infect humans in different parts of the world. It is a non-motile rod first discovered in 1882 by Robert Koch. It stains with difficulty but once stained it resists decolourization with weak acids and alcohols. Due to this property it is termed as acid fast bacillus (AFB). This property of alcohol fastness and acid fastness is due to its special cell wall. Resent techniques like nuclear magnetic resonance (NMR) and mass spectral analysis have resulted in thorough understanding of this structure along with interpretation of M. Tuberculosis genome. The wall as understood today comprises of a peptidoglycan covalently attached via a linker unit to a linear galactofuran which in turn is attached to several strands of highly branched arabinofurans. The arabinofurans are then attached to mycolic acids. Within this lipid environment provided by mycolic acid are other lipids like cord factor and sulphatides etc which are responsible for pathogenicity of this organism. Mycolic acids are alkyl β hydroxyl fatty acids with exceptional long chains. Chain lengths may vary but may be up to 80 carbon atoms. In M tuberculosis, they are characterized by very hydrophobic C34 to C33 fatty acids with C32 to C24 α side chains. Mycolic acid content of M. Tuberculosis determines its pathogenicity and drug resistance.

Due to this special cell wall M. tuberculosis has special staining properties. The do not take gram stain like most other bacteria. It was the work of Paul Ehrlich, Friedrich Neelsen and Franz Ziehl that a reliable method of staining was developed after discovery of this bacillus by Robert Koch. Staining of M. Tuberculosis in tissue is a great challenge and staining yield of this bacteria in tissue is very low. Nonetheless it is more important to demonstrate M. tuberculosis in granulomas to make a definitive diagnosis of tuberculosis because of its treatable nature. Importance of demonstration of M tuberculosis in tissue is further increased by the fact that TB granulomas may...
mimic leprosy, sarcoidosis and a number of fungal conditions.\textsuperscript{17}

**MATERIAL AND METHODS**

This was a cross-sectional comparative study. The study was conducted in Department of Pathology PGMI Lahore in collaboration with Pathology Department Ayub Medical College Abbottabad. Samples were collected from 27\textsuperscript{th} Dec 2011 to 29\textsuperscript{th} Jan 2014. It was a multi-faceted study and was concluded on 18\textsuperscript{th} April 2014. A total of 50 histopathologically diagnosed cases of tuberculous lymphadenopathy were included in the study. It was a non-probability purposive sampling. Patient history was taken and recorded on a proforma. Those patients with proven tuberculosis on histopathology and without any concomitant disease were selected. Tissues were routinely processed and wax embedded. Three-micron thick sections were taken and stained for H & E Zn and Auramin/ Rhodamin stain. H & E and Zn stained sections were studied under light microscope Olympus model CH. Auramin/ rhodamin stained section was studied under BHS-BH\textsubscript{2} Olympus with florescent attachments. Sections were studied under ultraviolet excitation frequency of 490° A.

Both Zn & Auramin /Rhodamin stained sections were visualized with 100X oil emersion lens in their respective microscopes. Presence of *M. tuberculosis* as red rods in ZN stain and Greenish yellow fluorescence in Auramin/Rhodamin stain was taken as positive for staining.

**RESULTS**

Out of the 50 diagnosed cases 17 were males and 33 were females (Figure-1). Patients had an age range of 9–80 (Figure-2) years with a mean age of 26–86±13 years. Eighteen presents (18\%) of the cases were positive for AFB on ZN Staining and 42\% were positive by Auramin/Rhodamin stain (Figure-3).

All the cases detected by ZN stain were picked up by Auramin/Rhodamin stain. Statistical analysis was done by using 2×2 table. Results of two stains were compared and calculated and was seen that Auramin/Rhodamin stain was significantly better than ZN stain. The *p*-value calculated was 0.0008 was less than .05 hence was significant.

As we have two rows and two columns, calculated degree of freedom is 1. Taking *p* value at less than 0.05 the expected chi square value should be above 3.84. Our calculated value is 6.88 giving a *p* value of 0.00088. This value is less than 0.05 hence highly significant. It proves that Auramine/Rhodamine is a better stain then ZN stain for tissue staining of *M. Tuberculosis*.

![Figure-1: Sex of the patient](image1)

![Figure-2: Age of the patients](image2)

![Figure-3: Stain positivity](image3)
DISCUSSION

Tuberculosis is an important disease of past and modern times.¹⁸ It is an important problem of developing countries and becomes more important when coexists with AIDS.¹⁹ Extra pulmonary TB is common in Pakistan. Among other diagnostic modalities biopsy is a slandered procedure and test. For definitive diagnosis, it is essential to demonstrate M. tuberculosis in tissue sections and tuberculous lesion (granulomas) because of very large differential for tuberculosis.¹⁷ Staining of AFB in tissue is difficult and positive yield is low.¹⁶ To overcome this difficulty different stains have been tried with variable success. Current study focuses on Routine ZN stain and Auramin/Rhodamin stain.

Results of our study shows that 86% of the patients were below the age of 40 years this in conformity with many previous studies as Muynck et al.²⁰ This also shows more trend of spread in younger people. Our study shows a higher preponderance in female (66%). This is in accordance with regional study conducted by Mukherjee et al.¹⁶ in India. However, the WHO report⁴ contradicts the findings of our study by narrating a lower prevalence of TB in females in Pakistan but admits that it may be underestimated because of local customs.

Our study shows a 18% positivity of AFB in ZN staining method. Pawal et al.²¹ have shown 22% positivity which is close to the findings of our study. Karimi et al.²² has shown that ZN stain is a low sensitivity low specificity stain and further narrated that results in different case series are variable ranging from zero to 40%. These results are consistent with the results of our study. Mukherjee et al.¹⁶ in his study shows a positivity rate of 44% which are higher in comparison to our study. This could be because of immunosuppression prevalent in India due to poverty and higher prevalence of AIDS in India but he himself admits that it varies between 23–65 in different age groups.

Our study shows 42% positivity for Auramine/Rhodamin stain. Results of our study are lower than Sherestha et al. who claims 71% positivity with Auramin/Rhodamin stain. It is remarkable that the cases he considered were all culture positive. His gross positivity disregarding culture was 36% close to the findings of our study. Ghenaat et al.²³ has claimed a positivity of 26% with Auramine/Rhodamin stain in his study. His sample size was 40 cases 20% less than our study this might be the cause of low yield in his study.

During the study period, I visited many centres of histopathology in the country and have observed that inspite of the importance of demonstration of AFB in granulomas, staining for the same is not a standard practice. This may be because of low sensitivity of AFB in ZN staining procedure and lack of florescent microscope facility or poisonous nature of Auramine/ Rhodamin stain⁴ that requires strict safety protocol.

AUTHORS' CONTRIBUTION

FA: Data collection, data analysis. OJ: Data Analysis, Discussion. AR: Staining procedure, Tissue processing.

REFERENCES