The combined role of Nth and MutY DNA glycosylases in

*Mycobacterium smegmatis*

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A dissertation submitted to the Faculty of Health Sciences, University of the Witwatersrand, Johannesburg, in fulfilment of the requirements for the degree of Master of Science in Medicine.
Declaration

I declare that this dissertation is my own, unaided work. It is being submitted for the degree of Masters of Science at the University of the Witwatersrand, Johannesburg. It has not been submitted before for any degree or examination at any other University.

__________________________________________________________________________

(Gadisi Nthambeleni) _______________ day of ___________ 2016
Abstract

During infection, *Mycobacterium tuberculosis* (MTB) encounters hostile conditions such as nutrient starvation, hypoxia and low pH which results in the generation of host-derived reactive oxygen (ROS) and nitrogen species (RNS) as part of the immune response to control the infection. Exposure to these reactive radicals can lead to oxidative damage of DNA which ultimately creates genomic instability through the introduction of mutations. Mycobacterial species have a high G+C content and these bases are particularly prone to oxidative damage. To counter this, MTB possesses specialized DNA repair systems such as the multi-step, multi-enzyme base excision repair (BER) pathway, wherein repair of oxidative damage is initiated by DNA glycosylases. These DNA glycosylases include the endonuclease III (Nth), endonuclease VIII (Nei) and formamidopyrimidine glycosylases (Fpg). In addition, MutY which acts together with Fpg (MutM) and MutT, forms part of the GO system, preventing mutations produced from 7,8-dihydro-8-oxoguanine (8-oxoG) lesions. Previously in our laboratory, we demonstrated a novel antimutator role for Nth and MutY DNA glycosylases in the BER pathway for the maintenance of mycobacterial genome stability. These data showed an increase in spontaneous mutation rate and decreased survival of *Mycobacterium smegmatis* when the *mutY* gene was disrupted together with the *fpg* DNA glycosylases. Additionally, deletion of the *nth* DNA glycosylase in *M. smegmatis* resulted in increased mutation rates under DNA damaging conditions and its inactivation in combination with *nei* resulted in reduced survival in an oxidative environment, with a heightened mutation frequency. In this study, we further investigated the combined role of Nth and MutY in an attempt to uncover the molecular mechanisms that protect mycobacterial DNA under hostile host environments. Double deletion mutants of *M. smegmatis* mc²155 lacking both the *mutY* and *nth* genes (ΔmutYΔnth, ΔnthΔmutY) were generated by homologous recombination. Both mutants displayed no growth defects under normal culture conditions (7H9 media) when
compared to the wildtype mc\(^2\)155 or the respective single \(M.\ smearatis \Delta nth\) and \(\Delta mutY\) mutants. Under \textit{in vitro} oxidative stress conditions, as generated by hydrogen peroxide, both the double deletion mutants displayed reduced survival kinetics compared to mc\(^2\)155 after 6 hours of exposure to hydrogen peroxide. As previously observed, loss of the \(nth\) gene resulted in increased DNA damage-induced mutation frequencies to rifampicin. In contrast, the \(mutY\) single deletion mutant and the double deletion mutants lacking both \(nth\) and \(mutY\) did not show increased DNA damage-induced mutagenesis compared to mc\(^2\)155. However, in the fluctuation assay, both double mutant strains, \(\Delta nth\Delta mutY\) and \(\Delta mutY\Delta nth\), displayed an increase in spontaneous mutation rates to rifampicin when compared to wild type and the single \(\Delta nth/\Delta mutY\) deletion mutants. The \(\Delta nth\Delta mutY\) mutant demonstrated an exacerbated mutation rate when compared to the \(\Delta mutY\Delta nth\) mutant. Collectively, these data reinforce the previously observed antimutator role for the mycobacterial MutY and Nth DNA glycosylases. The exacerbated phenotype observed for \(\Delta nth\Delta mutY\) suggest that there is a functional hierarchy between the various DNA glycosylases in the BER pathway, with the Nth DNA glycosylase superseding MutY as an antimutator in mycobacterial genome maintenance.
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Presentations

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<tr>
<td>A</td>
<td>Adenine</td>
</tr>
<tr>
<td>AIDS</td>
<td>Acquired Immune Deficiency Syndrome</td>
</tr>
<tr>
<td>Amp</td>
<td>Ampicillin</td>
</tr>
<tr>
<td>AP</td>
<td>Apurinic/Apyrimidinic</td>
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<tr>
<td>aph</td>
<td>Gene Encoding Kanamycin Resistance</td>
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<tr>
<td>BCG</td>
<td>Bacille Calmette-Guerin</td>
</tr>
<tr>
<td>BER</td>
<td>Base Excision Repair</td>
</tr>
<tr>
<td>BLAST</td>
<td>Basic Local Alignment Search Tool</td>
</tr>
<tr>
<td>bp</td>
<td>Base Pairs</td>
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<tr>
<td>BSA</td>
<td>Bovine Serum Albumin</td>
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<tr>
<td>C</td>
<td>Cytosine</td>
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<tr>
<td>CFU</td>
<td>Colony Forming Unit</td>
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<td>CFZ</td>
<td>Clofazimine</td>
</tr>
<tr>
<td>CTAB</td>
<td>Cetyltrimethylammonium Bromide</td>
</tr>
<tr>
<td>DCO</td>
<td>Double Cross Over</td>
</tr>
<tr>
<td>DIG</td>
<td>Digoxigenin</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethylsulphoxide</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic Acid</td>
</tr>
<tr>
<td>DNase</td>
<td>Deoxyribonuclease</td>
</tr>
<tr>
<td>DNTPs</td>
<td>Deoxynucleotide Triphosphate</td>
</tr>
<tr>
<td>DOTS</td>
<td>Directly Observed Treatment Short Course</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<tr>
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<tr>
<td>DS</td>
<td>Downstream</td>
</tr>
<tr>
<td>EDTA</td>
<td>Ethylene Diamine Tetraacetic Acid</td>
</tr>
<tr>
<td>EMB</td>
<td>Ethambutol</td>
</tr>
<tr>
<td>EtOH</td>
<td>Ethanol</td>
</tr>
<tr>
<td>Fpg</td>
<td>Formamidopyrimidine DNA Glycosylase</td>
</tr>
<tr>
<td>g</td>
<td>Gram</td>
</tr>
<tr>
<td>G</td>
<td>Guanine</td>
</tr>
<tr>
<td>GFX</td>
<td>Gatifloxacin</td>
</tr>
<tr>
<td>G+C</td>
<td>Guanine and Cytosine</td>
</tr>
<tr>
<td>HIV</td>
<td>Human Immunodeficiency Virus</td>
</tr>
<tr>
<td>INH</td>
<td>Isoniazid</td>
</tr>
<tr>
<td>iNOS</td>
<td>Inducible Nitric Oxide Synthase</td>
</tr>
<tr>
<td>kan</td>
<td>Kanamycin</td>
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<tr>
<td>LA</td>
<td>Luria-Bertani Agar</td>
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<tr>
<td>lacZ</td>
<td>Gene Encoding β-galactosidase</td>
</tr>
<tr>
<td>LB</td>
<td>Luria-Bertani Broth</td>
</tr>
<tr>
<td>LTBI</td>
<td>Latent Tuberculosis Infection</td>
</tr>
<tr>
<td>m</td>
<td>Mili</td>
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<tr>
<td>M</td>
<td>Molar</td>
</tr>
<tr>
<td>MDR-TB</td>
<td>Multidrug-Resistant Tuberculosis</td>
</tr>
<tr>
<td>ml</td>
<td>Millilitre</td>
</tr>
<tr>
<td>MMR</td>
<td>Mismatch Repair</td>
</tr>
<tr>
<td>Min</td>
<td>Minutes</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Definition</td>
</tr>
<tr>
<td>--------------</td>
<td>------------</td>
</tr>
<tr>
<td>MutY</td>
<td>Adenine DNA Glycosylase</td>
</tr>
<tr>
<td>m-value</td>
<td>Number of Mutational Events</td>
</tr>
<tr>
<td>n</td>
<td>Nano</td>
</tr>
<tr>
<td>NaCl</td>
<td>Sodium Chloride</td>
</tr>
<tr>
<td>NaOH</td>
<td>Sodium Hydroxide</td>
</tr>
<tr>
<td>Nei</td>
<td>Endonuclease VIII</td>
</tr>
<tr>
<td>NER</td>
<td>Nucleotide Excision Repair</td>
</tr>
<tr>
<td>NOD</td>
<td>Nucleotide –Binding Oligmerazation Domain</td>
</tr>
<tr>
<td>Nth</td>
<td>Endonuclease III</td>
</tr>
<tr>
<td>OD&lt;sub&gt;600nm&lt;/sub&gt;</td>
<td>Optical Density at Wavelength of 600 nm</td>
</tr>
<tr>
<td>PAMP</td>
<td>Pathogen Associated Molecular Patterns</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase Chain Reaction</td>
</tr>
<tr>
<td>PRR</td>
<td>Pattern Recognition Receptors</td>
</tr>
<tr>
<td>PTH</td>
<td>Prothionamide</td>
</tr>
<tr>
<td>PZA</td>
<td>Pyrazinamide</td>
</tr>
<tr>
<td>rif&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Rifampicin Resistance</td>
</tr>
<tr>
<td>Rpm</td>
<td>Revolutions per minute</td>
</tr>
<tr>
<td>RRDR</td>
<td>Rifampicin Resistance Determining Region</td>
</tr>
<tr>
<td>SCO</td>
<td>Single Cross Over</td>
</tr>
<tr>
<td>sdH&lt;sub&gt;2&lt;/sub&gt;O</td>
<td>Sterile Distilled Water</td>
</tr>
<tr>
<td>sac&lt;sub&gt;B&lt;/sub&gt;</td>
<td>Gene Encoding Levansucrase</td>
</tr>
<tr>
<td>SDS</td>
<td>Sodium Dodecyl Sulphate</td>
</tr>
<tr>
<td>T</td>
<td>Thymine</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>TB</td>
<td>Tuberculosis</td>
</tr>
<tr>
<td>TDR-TB</td>
<td>Total Drug Resistance TB</td>
</tr>
<tr>
<td>TNF</td>
<td>Tumour Necrosis Factors</td>
</tr>
<tr>
<td>TLR</td>
<td>Toll Like Receptors</td>
</tr>
<tr>
<td>Tris</td>
<td>Tris (hydroxymethyl) aminomethane</td>
</tr>
<tr>
<td>Tween</td>
<td>Polyoxyethylene Sorbitan Monooleate</td>
</tr>
<tr>
<td>US</td>
<td>Up Stream</td>
</tr>
<tr>
<td>UV</td>
<td>Ultraviolet Light</td>
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</table>
1. Introduction

1.1 Tuberculosis

The genus *Mycobacterium* contains many important pathogenic bacteria such as *Mycobacterium leprae*, *Mycobacterium ulcerans* and *Mycobacterium bovis* which cause infections in both humans and animals. In humans, *Mycobacterium tuberculosis* (MTB) is the causative agent of tuberculosis (TB) which primarily infects the respiratory system. Robert Koch discovered the link between MTB and TB in 1882. Mycobacterial species are representative of both Gram-negative and Gram-positives (more related to Gram-negative), rod shaped bacteria with a diameter of between 0.2 -0.6 µm and length of 2-5 µm (Fu and Fu-Liu, 2002, Stoops et al., 2010). However, the gram classification of mycobacteria is problematic as it does not consider the mycolic acid layer in mycobacterial cell wall, which acts an outer membrane but is not comprised of phospholipids.

TB accounts for an estimated 1.4 million deaths and 9.6 million new cases of infection annually (WHO, 2015). In 1993, the World Health Organization (WHO) declared TB a global health emergency (WHO, 1993). Sub-Saharan Africa has the highest prevalence of TB worldwide and it is estimated that 254 000 adults and children died from TB related cases per year (Keugoung et al., 2014, Zumla, 2015). Moreover, people with compromised immune systems are more susceptible to TB hence, this opportunistic pathogen has consistently been one of the main causes of death in people living with Human Immunodeficiency Virus (HIV) or Acquired Immunodeficiency Syndrome (AIDS) (Rana et al., 2000, Corbett et al., 2003,Karim et al., 2009), with 15 % of TB incident cases being those co-infected with HIV/AIDS (WHO, 2009). This situation is further worsened by the rapid emergence of multi-drug resistant (MDR) forms of TB strains, which have acquired resistance against commonly used first line drugs such as rifampicin and isoniazid, extensively-drug resistant (XDR)
strains that are MDR but have also developed resistance to fluroquinolones and aminoglycosides. Totally drug resistant (TDR) strains of MTB are resistant to all first line and second line drugs (Zignol et al., 2006, Andrews et al., 2007, Hall et al., 2009, Calver et al., 2010, Gandhi et al., 2010).

1.2 Tuberculosis Therapy

The only available and licensed vaccine for TB is M. bovis BCG (Bacille Clammete–Guerin), made from an attenuated strain of Mycobacterium bovis. BCG is usually given to new-born babies immediately after birth. However, the limitation of BCG is its varying efficacy against pulmonary tuberculosis in different settings, ranging from 60% to 80% in both infants and children (Colditz et al., 1994, Rodrigues et al., 1993, Fine, 1995). The reasons for this variation in efficacy include genetic variation in strains, genetic variation in human populations and geographical location (Brewer, 2000, Rowland and McShane, 2011).

The currently available first line TB drugs are more than 40 years old. Hence, there is a dire need for novel drugs for the treatment of TB, particularly for the treatment of MDR, XDR and TDR strains. The first line TB drug regimen includes a four-drug cocktail comprising of rifampicin, isoniazid, pyrazinamide and ethambutol for 6 months for the treatment of drug susceptible strains (McIlleron et al., 2006, Ma et al., 2010). As mentioned previously, the rapid emergence of drug resistant strains has rendered many of these treatments ineffective, thus requiring prolonged treatment of approximately 18 months and the use of second line drugs such as cycloserine, ethionamide, p-aminosalicylic acid (PAS), kanamycin and fluoroquinolones (Chhabra et al., 2012, Singh et al., 2012). This lengthy treatment creates two major challenges which includes, managing the toxicity of the drugs and making sure that the patient is compliant until the end of treatment (Longo et al., 2015). Countries, like
Bangladesh and Cameron, have shortened the time for the treatment of MDR-TB from 20 to 9 months with the use of a seven drug regimen, with a cure rate of 85 to 89% (Aung et al., 2014, Kuaban et al., 2015). The 9 month regimen includes high dosage of ethambutol (EMB), gatifloxacin (GFX), clofazimine (CFZ), pyrazinamide (PZA), prothionamide (PTH) followed by a 4 month intensive phase with, kanamycin (kan) and isoniazid (INH) (Aung et al., 2014).

Directly Observed Treatment Short Course (DOTS) is a highly recommended approach by WHO for fighting TB. It involves observation of the patient for 6 to 18 months by a health care worker to ensure that the patient swallows each of the anti-TB drugs prescribed, at each dosing interval (Mitchison, 1985, Sagbakken et al., 2013). The main goals of DOTS are to improve the cure rate, to monitor patient compliance and to prevent relapses due to MDR-TB (Cox et al., 2006, Frieden and Sbarbaro, 2007).

1.3 Tuberculosis infection

The lung alveolar macrophages are the first line of defence against TB infection and phagocytose the bacilli. Monocytes and neutrophils are also recruited to the site of infection to assist with clearance of the pathogen. The membrane bound pattern recognition receptors (PRR)s such as Toll-like receptors (TLRs) and C-type lectins (TLRs) or cytoplasmic receptors such as the nucleotide-binding oligomerization domain-(NOD) containing proteins recognize the pathogen via the pathogen-associated molecular patterns (PAMPs) (Akira et al., 2006, Harding and Boom, 2010). The interaction between PPR and PAMPs triggers the innate immune response, leading to mild inflammation, initiating the antimicrobial response (Mogensen, 2009, Ahmad, 2010). Antimicrobial responses activate the transcription factor (NF)-κB and the production of cytokines such as tumor necrosis factor (TNF)-α, interleukin
IL-1, IL-12, chemokines and nitric oxide (Caamano and Hunter, 2002, Ahmad, 2010). An important mechanism that enables MTB to survive in macrophages is the ability to prevent maturation of the phagosomes into phagolysosomes, which allow the bacteria to multiply in the phagosome (Ahmad, 2010). In addition, T-cell cytokines, chemokines and other host cells, such as dendritic cells and endothelial cells, are also deployed to the site of infection as part of the adaptive immune response (Sasindran and Torrelles, 2011, Silva Miranda et al., 2012). This results in the formation of granulomas, widely considered a hallmark of tuberculosis infection (Sasindran and Torrelles, 2011, Silva Miranda et al., 2012). Granulomas are a collection of immune cells which cluster around the bacteria, acting as a blockade, effectively restricting diseases spread to other parts of the lung and are clearly visible on a chest X-ray, serving as a good indication of TB disease (Woodworth and Behar, 2006, Ramakrishnan, 2012). However, during infection bacilli are able to escape this harsh immune assault generated by the host immune system by transitioning into a state of low metabolic activity, also known as bacterial persistence (Flynn and Chan, 2001, Gengenbacher and Kaufmann, 2012).

Latent TB infection which can be diagnosed by a tuberculin skin test, describes an individual who is infected with TB but displays no symptoms of TB disease and is consequently not contagious (Barry et al., 2009, Chao and Rubin, 2010). It is estimated that more than one third of the humanity is latently infected with MTB and these individuals serve as a large reservoir for future active disease as they carry a 10% annual risk of developing active TB through the process of reactivation (Dooley Jr et al., 1990, Corbett et al., 2003). Risk factors that contribute to reactivation of latent TB range from HIV co-infection, host genetics, malnutrition, indoor air pollution, smoking or diabetes (Jick et al., 2006, Lönnroth and Raviglione, 2008). Studies have shown that people infected with HIV have a higher risk of reactivation compared to non-infected individuals (Horsburgh Jr, 2004).
Another efficient method used by infected macrophages to control infection is the production of reactive oxygen species (ROS) such as hydroxyl radicals (•OH), singlet oxygen (¹O₂), hydrogen peroxide (H₂O₂) and reactive nitrogen species (RNS) (Zahrt and Deretic, 2002, Cadet and Wagner, 2014). ROS and RNS have the ability to cause damage to proteins, lipids and nucleotides (DNA), thus leading to either cell death or genomic instability. ROS and RNS are produced in activated macrophages by two enzymes namely NADPH oxidase (NOX2/gp91phox) and inducible nitric oxide synthase (iNOS) respectively (Ehrt and Schnappinger, 2009). The superoxide anions (O₂⁻) produced by NOX2 serve as a catalyst for the production of ROS (Babior, 1999). ROS are formed when NOX2 assembles into an active enzyme complex that transports electrons across the membrane from NADPH to molecular oxygen, which results in the production of O₂⁻, which dismutates into H₂O₂ and generates lethal hydroxyl radicals (Ehrt and Schnappinger, 2009). IFNγ induces iNOS and generates nitrite along with nitrate through metabolism of nitric oxide.

1.4 DNA damage caused by oxidative stress

As mentioned previously, ROS and RNS also have the ability to inflict damage on DNA, and such damage often results in cell death or alternatively in genetic instability leading to mutations. The hydroxyl radical (•OH) is one of the most reactive free radicals, resulting in damage to DNA such as single or double stranded breaks, mismatch base pairing, as well as apurinic and apyrimidinic (AP) sites (Demple and Harrison, 1994, Dos Vultos et al., 2009, Bridge et al., 2014). Mycobacteria are mostly susceptible to oxidative damage because of the high G+C content in their genomes (Dos Vultos et al., 2009, van der Veen and Tang, 2015). Guanine is most vulnerable to oxidative DNA damage when compared to other bases due to its high oxidation potential, resulting in lesions such as 8-oxo-7,8-dihydroguanine (8-oxoG), which is pro-mutagenic and 2,6-diamino-4-hydroxy-5-formamido-pyrimidine (FapyG),
which is cytotoxic as it has the ability to block replication (Laval et al., 1998, David et al., 2007). The 8-oxoG lesion may result in G:C → T:A transversions, resulting in mispairing with adenine during replication if not repaired (Maki and Sekiguchi, 1992, Hazra et al., 2001). 8-oxoG can be further oxidized to spirominodihydrantoin and guanidinohydantoin (Luo et al., 2000, Hailer et al., 2005). Other base adducts caused by these hydroxyl radicals and free radicals are oxidation of cytosine to hypoxanthine, formed as a result of oxidation of the 5,6 double bond of cytosine, producing cytosine glycol (von Sonntag and Schuchmann, 1994, Cadet et al., 1999, Gros et al., 2002). Hydroxyl radical attacks on adenine generate oxidative adenine derived products such as 4, 6-diamino-5-formamidopyrimidine (FapyAde) and 8-oxo-7, 8-dihydroadenine (8-oxoAde), resulting in AT → GC mutations (Tudek, 2003). When thymine is exposed to oxidative stress, it results in changes in molecular structure and produces substrates such as 4,5 dihydroxyhydrothymine know as thymine glycol, 5-hydroxymethyluracil (hmU) and 5-formyluracil (fU), which is one of the major oxidative thymine lesions (Masaoka et al., 2001, Pfaffeneder et al., 2014).

1.5 Mycobacterium tuberculosis defence against oxidative stress

To defend themselves against damage caused by oxidative stress, pathogens have evolved several mechanisms responsible for resisting ROS and RNS, which involve the use of detoxifying molecules, the scavenging of ROS by certain vitamins and enzymes or protein regulation (Asad et al., 2004, Gutteridge and Halliwell, 2010). Detoxifying mechanisms against oxidative stress have been well characterized in Escherichia coli (E. coli) (Storz and Imlayt, 1999). When E. coli is exposed to oxidative stress, OxyR and SoxRS induces the expression of oxidative stress response genes (Zheng et al., 1998). OxyR stimulates the expression of a set of antioxidant genes which includes the catalases katG, katE and katF,
**ahpCF** (alkyl hydroperoxidase) and **oxyS** (a regulatory RNA) (Zheng et al., 1998, Zheng et al., 2001). The SoxRS regulon also triggered the expression of **sodA** and **sodB** which are superoxide dismutases (Touati, 1988, Bébien et al., 2002). Through redox-responsive 2Fe-2S clusters, SoxRS controls the expression of oxidative stress responsive genes by sensing nitrogen and oxidative stress (Green and Paget, 2004).

Interestingly in mycobacterial species the homologue for SoxR is missing, while OxyR is non-functional (Deretic et al., 1995, Chawla et al., 2012). Studies have proposed that in order to manage ROS/RNS stress during infection, mycobacteria sense redox changes and make use of the heme-based DosR/S/T system along with the thiol-based SigH/RshA system as a redox sensor (den Hengst and Buttner, 2008). In addition, the KatG catalase-peroxidase is a fascinating oxidative protective enzyme, which efficiently and effectively decomposes hydrogen peroxide (H$_2$O$_2$) into water and oxygen (Ng et al., 2004, Vlasits et al., 2007). Other important enzymatic defensive mechanisms against these intermediates are alkyl hydroperoxide reductases such as AhpC which scavenges H$_2$O$_2$ from the host (Chen et al., 1998, Springer et al., 2001).

### 1.6 DNA repair systems

DNA integrity is important for optimal cellular functioning and bioinformatics analysis of the sequenced MTB genome revealed the presence of homologs involved in several DNA repair pathways, namely nucleotide excision repair (NER), base excision repair (BER), SOS and recombination repair, which together detect and repair DNA damage to prevent cell lysis and mutation (Mizrahi and Andersen, 1998, Cole et al., 1998, Güthlein et al., 2009). Notably, in mycobacteria an important repair pathway is absent, the DNA mismatch repair system which may have serious repercussions (causing genome instability) for the repair system of this
genus (Mizrahi and Andersen, 1998). However, despite this the MTB genome is stable, suggesting that it is able to repair mispaired bases using other repair systems.

1.6.1 SOS repair system

The essential components of the SOS repair system are the LexA (dimer) and RecA proteins, which act as a repressor and inducer respectively by stalling DNA replication (Brent and Ptashne, 1981, Boutry et al., 2013). When there is no DNA damage LexA binds to a specific sequence known as the SOS box, upstream of the recA and lexA genes and represses their expression (Little et al., 1981, Davis et al., 2002). When DNA damage occurs, RecA coordinates and stimulates the expression of the genes required for survival by binding to the single-stranded DNA, resulting in the formation of a nucleoprotein filament. This action stimulates auto-cleavage, with LexA activating the SOS genes involved in recombination repair (Davis et al., 2002, Žgur-Bertok, 2013). Bioinformatics analysis has demonstrated that MTB possesses all the required genes needed for a functional SOS system (Cole et al., 1998, Mizrahi and Andersen, 1998). The main role of the SOS repair system is to induce genes involved in DNA repair, including base excision repair (BER), nucleotide excision repair (NER) and recombination DNA repair depending on which is suitable for the type of damage (Friedberg et al., 2005). All the known mycobacterial DNA repair systems will be discussed further below.

1.6.2 Recombination repair

Recombination is an important biological process wherein two DNA molecules combine or exchange genetic material. This process can be either homologous (exchange of DNA
molecules occurring between two identical molecules) or non-homologous (exchange of DNA molecules without sequence homology). Double stranded breaks caused by free radicals, radiation or replication errors are normally fixed by homologous recombination repair (Kuzminov, 1999). In mycobacteria and other bacteria, the main mediator of recombination repair system is the recA gene (Muttucumaru and Parish, 2004). Briefly, when a double stranded break is recognised in the genome, it causes RecB, RecC and RecD to form a three subunit complex, RecBCD, which is bifunctional with both helicase and nuclease activity (Dillingham et al., 2003). Once this complex attaches to the ends, it rapidly digests the DNA in a 3′ → 5′ fashion. RecA assists specifically in pairing the single stranded DNA molecule with a homologous single stranded protein to repair the double stranded break (Dixon and Kowalczykowski, 1995, Dillingham et al., 2003). Also in MTB RecA contains an intein that has to spliced out for the protein to be functional (Davis et al., 2001, Sander et al., 2001). In other pathogens such as Salmonella typhimurium, a recA deficient mutant showed enhanced sensitivity to the DNA damaging agent, hydrogen peroxide (H$_2$O$_2$) (Buchmeier et al., 1995) indicating that RecA plays a significant role during DNA repair.

1.6.3 Nucleotide excision repair (NER)

One of the key DNA repair pathways is nucleotide excision repair (NER), which involves the excision of oligonucleotides that are approximately 30 base pairs long. The enzymes of the NER pathway recognize and eliminate bulky pyrimidine dimer lesions and 6,4 photoproducts caused by UV irradiation and genotoxic compounds (Van Houten et al., 2002, Rastogi et al., 2010). There are three main enzymes involved in the NER pathway in bacteria, these are UvrA, UvrB and UvrC (ABC complex), which are assisted by DNA polymerase I and DNA helicase II (UvrD) (Dos Vultos et al., 2009, Farnell, 2011). The NER pathway is a four step
system, which starts with the scanning and recognition of DNA lesions by protein trimmers, including two molecules of UvrA and UvrB (UvrAB machinery), followed by the release of the two UvrA molecules and pre-incision by UvrB, which splits the two DNA strands to validate the position of the lesion. UvrC binds to the site of the strand to be cleaved on the phosphodiester bond, and generates cuts several bases (4-5) away from the damaged site. The helicase UvrD, binds and displaces the segment of the damaged strand from the DNA molecule, followed by filling of the gap with the correct nucleotide by DNA polymerase I and sealing by DNA ligase (Sancar, 1996, Van Houten et al., 2002, Shuck et al., 2008, Kisker et al., 2013).

Interestingly, the mechanism of NER between eukaryotes and prokaryotes is similar however, there is a difference in the number of proteins involved, with more proteins required in eukaryotes compared to prokaryotes (Farnell, 2011). The main component of NER in bacteria is UvrB, which is responsible for damage recognition with two molecules of UvrA (Skorvaga et al., 2002). In E. coli K12, a uvrB mutant was more susceptible to psoralen (a photoactive chemical agent) when compared with uvrA and uvrC gene knockout mutants, indicative of a distinct role in the repair of adducts (Lage et al., 2010). In MTB, the UvrB mutant is highly sensitive to acid nitrite, used as a source of NOS (Darwin and Nathan, 2005). Additionally, the UvrB mutant of Mycobacterium smegmatis has increased sensitivity to UV, low pH, reactive oxygen species (ROS) and hypoxia when compared with the wildtype (Kurthkoti et al., 2008). Taken together, NER in mycobacteria appears to play an important role in protecting the bacterium from damage inflicted by the host immune system.
1.6.4 Base excision repair

Base excision repair (BER) is an important DNA repair pathway that deals mainly with oxidatively damaged bases (Lu et al., 2001, Kurthkoti and Varshney, 2011). DNA glycosylases are the first enzymes in the pathway that are responsible for the recognition and removal of damaged DNA, Figure 1 (Fromme et al., 2004, Guo et al., 2010, Jacobs and Schär, 2012). BER can be viewed in five basic steps shown in Figure 1: (I) the DNA glycosylase identifies and removes the oxidatively damaged or incorrect bases (shown in red), which results in (II) the formation of an AP (purinic/apyrimidinic site) site. (III), the AP endonuclease cleaves the sugar-phosphate backbone, creating a single stranded break, resulting in a 3′ hydroxyl group and 5′-terminal deoxyribosephosphate residue.(IV) deoxyribo phosphodiesterase (dRpase) excises the 5′-terminal deoxyribose phosphate and converts the 3′ termini to a hydroxyl group. (V) DNA polymerase then adds in the new correct nucleotide (shown in yellow) and DNA ligase seals the nick.

![Figure 1](image-url)  
**Figure 1.** The multi-step, multi enzyme base excision repair (BER) pathway showing the various steps involved in the repair of damaged bases. Modified from Kurthkoti and Varshney, 2011.
1.7 Glycosylase families involved in the base excision (BER) pathway

Glycosylases can be classified into families based on both sequence and structural homology. Even though structural and sequence homology may exists among members of the same family, these glycosylases can exhibit unique features and functions. In mycobacteria these glycosylases include the Endonuclease III (Nth) superfamily the formamidopyrimidine (Fpg/MutM/Fapy) and endonuclease VIII (Nei) DNA glycosylases which together form the Fpg/Nei family (Mizrahi and Andersen, 1998, Moolla et al., 2014, Hassim et al., 2015).

Fpg and Nei glycosylases are categorized as the Fpg/Nei family as the sequence of these enzymes shares some common structural and biochemical features such as a helix-two-turn-helix motif (H2TH) and a zinc finger motif (Zharkov et al., 2003, Prakash et al., 2012). Additionally, it contains N- and C-terminal domains connected by a flexible hinge (Zharkov et al., 2003). The N-terminal is composed of a β-sheet, whereas the α-helical C-terminal domain has a four cysteine zinc finger structure consisting of two anti-parallel β-strands, which are thought to be involved in DNA binding (Tchou et al., 1993, Zharkov et al., 2003). Both Fpg and Nei are bi-functional glycosylases as they possess both glycosylase and AP activity, which allows for the cleavage of the phosphodiester bond. Notably, substrate preferences of Fpg and Nei are different; the main substrate for Fpg is oxidized purines such as highly mutagenic 8-oxoG, while Nei recognizes oxidized pyrimidines such as the thymine intermediate dihydrothymine (DHT) (Wallace et al., 2003, Zharkov et al., 2003, Prakash et al., 2012). In MTB and the non-pathogenic faster growing close relative of MTB, Mycobacterium smegmatis (M. smegmatis), two copies of Fpg (FpgI and FpgII) and Nei (NeiI and NeiII) exist (Mizrahi and Andersen, 1998, Cole et al., 1998). However, in MTB FpgI is non-functional as it is truncated at the 3’ N-terminal and therefore, lacks the DNA binding domain required for glycosylase activity (Olsen et al., 2009, Guo et al., 2010).
The Nth and MutY DNA glycosylases are classified under the Nth super-family or Helix-hairpin-Helix (HhH) super-family (Thayer et al., 1995, Denver et al., 2003). Phylogenetic analyses indicates that the Nth protein is highly conserved in all phyla, with MutY found in only half of the Eukaryotes and less than half of the Archaeal species (Denver et al., 2003, Jacobs and Schär, 2012). Sequence alignments of the *E.coli* Nth and MutY glycosylases indicate that they share some common features such as the Helix-turn Helix-hairpin motif involved in DNA binding, a glycine-proline-aspartate (GPD) motif and the 4Fe-4S cluster loop (FCL), which is not involved in either the binding or removal of damaged bases (Thayer et al., 1995, Zharkov and Grollman, 2002, Markkanen et al., 2013). In terms of activity the Nth DNA glycosylase is classified as a bi-functional enzyme as it possesses both glycosylase and AP activity that allows for the cleavage of the phosphodiester bond in the damaged DNA (Lomax et al., 2005, Hegde et al., 2008). The classification of MutY is controversial with some studies classifying it as mono-functional and others as bi-functional, with weak 3’ AP lyase activity (Lu and Chang, 1988, Gogos et al., 1996, Corbett et al., 2003). Williams and David (1998) used an approach which traps the Schiff’s base intermediate between DNA glycosylases and their substrates and demonstrated that MutY is a mono-functional glycosylase which lacks AP lyase activity (Williams and David, 1998).

Although Nth and MutY are sequence related, they are functionally different and do not share the same primary substrate specificity (Zharkov and Grollman, 2002). The main substrate for the Nth is oxidized pyrimidines (Bauche and Laval, 1999, Zharkov and Grollman, 2002) whilst MutY recognizes adenine mispaired with 8-oxo-7,8-dihydroguanine (8-oxoG) (Michaels et al., 1990, Michaels and Miller, 1992). However, substrate overlap does exist between the Nth and Nei proteins as they share structural relatedness with Fpg and are able to recognize and repair oxidized purine lesions (Wallace, 1998, Prakash et al., 2012). In *Salmonella typhimurium*, mutation phenotypes only became noticeable with the
combinatorial deletion of the \textit{nth} and \textit{fpg/nei} genes, signifying an important antmutator role for \textit{nth} (Suvarnapunya et al., 2003). In support of this, a previous study conducted at the CBTBR has also demonstrated that lack of the Nth glycosylase in \textit{M. smegmatis} results in increased DNA damage-induced mutations when bacteria are exposed to UV light and the combinatorial loss of Nth and Nei resulted in reduced survival when challenged with oxidative stress (Moolla et al., 2014). The increase in mutation rate upon deletion of Nth, individually or in combination with Nei, suggests an important antmutator role for Nth as well as a novel interaction between the Nth and Nei DNA glycosylases (Moolla et al., 2014). In contrast, the combinational deletion of Nth and Fpg glycosylases did not result in increased mutation rates or reduced survival under oxidative stress (Moolla et al., 2014).

Organisms with G+C rich genomes like MTB are constantly at risk of being damaged by oxidative stress generated by the host, which may result in oxidative damage such as 8-oxoG (GO lesions) causing GC\textarrow{\rightarrow}TA and AT\textarrow{\rightarrow}CG mutations (Fraga et al., 1990, Steenken and Jovanovic, 1997). To counter such mutations, mycobacteria uses a DNA repair pathway known as the GO repair system, as shown in Figure 2 (Michaels and Miller, 1992, Kurthkoti and Varshney, 2012). The GO system involves the activity of three proteins, namely MutY, MutM (Fpg) and MutT, Figure 2 (Michaels and Miller, 1992, Tajiri et al., 1995). Briefly, (1) the guanine base is damaged by oxidative stress, forming an oxidized base 8-oxoG (red). (2) MutM/Fpg recognizes the damaged base pair, which is repaired by the BER pathway. (3) If replication takes place before the repair of the GO lesion by MutM/Fpg, this leads to the mis-incorporation of A with GO. (4) MutY binds to the mis-paired A against the GO lesion and offer a second chance to be repaired by Fpg. (5) The GO lesion is repaired by Fpg via the BER pathway. If repair is not achieved this results in a GC\textarrow{\rightarrow}AT transversion (red box) (Michaels and Miller, 1992, Kurthkoti and Varshney, 2012). MutT eliminates oxidized dGTP
from the nucleotide pool, preventing insertion of damaged guanine bases during replication thus avoiding mismatches with adenine (Sakumi et al., 1993, Dos Vultos et al., 2009).

Sanders et al. (2009) investigated the consequence of the lack of the GO pathway in the GC rich bacterium *Pseudomonas aeruginosa*, by inactivating *mutM, mutY* and *mutT* and found that this resulted in increased mutations under oxidative stress conditions. Kurthkoti et al. (2010) investigated the role of *mutY* in *M. smegmatis*, and found that the loss of *mutY* did not result in phenotypic changes or increased mutation rates under an oxidative environment (Kurthkoti et al., 2010). In contrast, recent studies in our laboratory demonstrated an increase in spontaneous mutation rates and decreased survival of *M. smegmatis* when the *mutY* gene was disrupted together with the *fpg/nei* DNA glycosylases, suggestive of an antimutator role for MutY (Hassim et al., 2015).

**Figure 2.** The GO pathway. Adapted from (de Oliveira et al., 2014).

**1.8 Aim and objectives**

The two previous studies at the CBTBR clearly demonstrated that the Nth and MutY DNA glycosylases are key enzymes of the BER pathway, responsible for preventing and or
reducing mutations in *M. smegmatis*. Hence, it is important to assess the combined role of these genes in maintaining mycobacterial genome integrity. In this study we aimed to assess the combinatorial role of these enzymes in BER. We hypothesize that the combined deletion of *nth* and *mutY* will lead to exaggerated mutation rates in *M. smegmatis*.

### 1.8.1 Specific objectives

Considering the above aims, the specific objectives were as follows:

- Generation of a *M. smegmatis* double deletion mutant strain by deleting the *mutY* gene in a Δ*nth* deletion mutant background and by deleting *nth* gene in a Δ*mutY*, using previously generated single deletion mutant strains and suicide vectors carrying inactivated copies of the *mutY* and *nth* genes respectively (Moolla *et al.*, 2014, Hassim *et al.*, 2015), Figure 3.

![Figure 3](image)

**Figure 3.** Strategy for generating combinatorial deletion mutants of Nth and MutY DNA glycosylases in *M. smegmatis*

- Genotypic verification of the mutant and parental strains using PCR and Southern blot analysis.

- Complementation of mutant strains by reintroducing the respective functional gene into the mutant strain to confirm restoration of the observed phenotypes, Figure 3 red.
• Phenotypic characterization of the *M. smegmatis* mc²155, single, double mutant and complemented strains by comparing the respective strains for:
  o Growth kinetics under normal growth conditions
  o Survival of the mutant and parental strains under oxidative stress conditions as generated by hydrogen peroxide.
  o Changes to rifampicin resistance.
  o Changes in spontaneous mutation rates as measured by the fluctuation assay.
  o Spectral assessment of the rifampicin resistant colonies isolated from the fluctuation assay to determine the genotype of the mutation that results in rifampicin resistance.

2. Materials and Methods

2.1 Bacterial strains and culture conditions

*E. coli* strains were grown overnight in a shaking incubator (Labcon shaking incubator) at 37°C in Luria-Bertani broth (1% tryptone, 1% NaCl, 0.5% yeast extract) (LB). LB media was supplemented with kanamycin (kan) at 50 µg/ml. *E. coli* strains transformed with large plasmids (>8kb) were grown at 30°C with shaking for two nights to prevent plasmid rearrangements. *M. smegmatis* and derivative strains were grown in Middlebrook 7H9 liquid medium (Difco) supplemented with 0.2% glycerol and 0.05% Tween80 (referred to hereafter as 7H9) with shaking or on Middlebrook 7H10 solid medium (Difco) supplemented with 0.085% NaCl, 0.2% glucose and 0.5% glycerol (referred to hereafter as 7H10). *M. smegmatis* complemented strains were grown in 7H9 supplemented with 25 µg/ml of kan.
### Table 2.1. Bacterial strains used in this study.

<table>
<thead>
<tr>
<th>Bacterial Strain</th>
<th>Characteristics</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>mc²155</td>
<td>High frequency transformation mutant of <em>M. smegmatis</em> ATCC 607</td>
<td>(Snapper <em>et al.</em>, 1990)</td>
</tr>
<tr>
<td>Δnth</td>
<td>Derivative of <em>M. smegmatis</em> mc²155 carrying an internal 570 bp deletion in <em>nth</em></td>
<td>(Moolla <em>et al.</em>, 2014)</td>
</tr>
<tr>
<td>ΔmutY</td>
<td>Derivative of <em>M. smegmatis</em> mc²155 containing an internal 682 bp deletion in <em>mutY</em></td>
<td>(Hassim <em>et al.</em>, 2015)</td>
</tr>
<tr>
<td>ΔnthΔmutY</td>
<td>Derivative of Δnth carrying an internal 682 bp deletion in <em>mutY</em></td>
<td>This study</td>
</tr>
<tr>
<td>ΔmutYΔnth</td>
<td>Derivative of ΔmutY carrying and internal 570 bp deletion in Δnth</td>
<td>This study</td>
</tr>
<tr>
<td>Δnth::nth</td>
<td>Derivative of Δnth carrying pTWEETY::nth integrated at the <em>attP</em> phage attachment site</td>
<td>(Moolla <em>et al.</em>, 2014)</td>
</tr>
<tr>
<td>ΔmutY::mutY</td>
<td>Derivative of ΔmutY carrying pTWEETY::mutY integrated at the <em>attP</em> phage attachment site</td>
<td>(Hassim <em>et al.</em>, 2015)</td>
</tr>
<tr>
<td>ΔnthΔmutY::mutY</td>
<td>Derivative of ΔnthΔmutY carrying pTWEETY::mutY integrated at the <em>attP</em> phage attachment site</td>
<td>This study</td>
</tr>
<tr>
<td>ΔmutYΔnth::nth</td>
<td>Derivative ΔmutYΔnth carrying pTWEETY::nth integrated at the <em>attP</em> phage attachment site</td>
<td>This study</td>
</tr>
<tr>
<td>DH5α</td>
<td><em>supE44 ΔlacU169 hsdR17 recA1 endA1 gyrA96 thi-1 relA1</em></td>
<td>(Hanahan, 1983)</td>
</tr>
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</table>

### Table 2.2. Vectors used previously and in this study. Plasmid maps are shown in Appendix 5.3.
<table>
<thead>
<tr>
<th>Plasmids</th>
<th>Characteristics</th>
<th>Size(bp)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>pGEM3Zf(+)</td>
<td>E. coli cloning vector; amp&lt;sup&gt;R&lt;/sup&gt;; lacZ-alpha; oriE</td>
<td>3199</td>
<td>Promega</td>
</tr>
<tr>
<td>p2NIL</td>
<td>E. coli cloning vector and mycobacterial suicide plasmid; kan&lt;sup&gt;R&lt;/sup&gt;; oriE</td>
<td>4753</td>
<td>(Parish and Stoker, 2000)</td>
</tr>
<tr>
<td>pGOAL17</td>
<td>Plasmid carrying lacZ and sacB genes as a PacI cassette; amp&lt;sup&gt;R&lt;/sup&gt;; oriE</td>
<td>8855</td>
<td>(Parish and Stoker, 2000)</td>
</tr>
<tr>
<td>pGOAL19</td>
<td>Plasmid carrying lacZ, hyg and sacB genes as a PacI cassette; amp&lt;sup&gt;R&lt;/sup&gt;, oriE</td>
<td>10435</td>
<td>(Parish and Stoker, 2000)</td>
</tr>
<tr>
<td>p2NIL&lt;sub&gt;mutY&lt;/sub&gt;::pGOAL17</td>
<td>p2NIL suicide vector carrying the deleted mutY gene and the selectable and counter selectable markers (lacZ-sacB) from pGOAL17, kan&lt;sup&gt;R&lt;/sup&gt;; lacZ; sacB; oriE.</td>
<td>14508</td>
<td>(Hassim et al., 2015)</td>
</tr>
<tr>
<td>p2NIL&lt;sub&gt;nth&lt;/sub&gt;::pGOAL19</td>
<td>p2NIL suicide vector carrying the deleted nth gene and the selectable and counter selectable markers (hyg-lacZ-sacB) from pGOAL19; kan&lt;sup&gt;R&lt;/sup&gt;hyg&lt;sup&gt;R&lt;/sup&gt;; lacZ; sacB; oriE.</td>
<td>14748</td>
<td>(Moolla et al., 2014)</td>
</tr>
<tr>
<td>pTWEETY</td>
<td>Mycobacterial integrating vector; kan&lt;sup&gt;R&lt;/sup&gt;, oriE, int</td>
<td>5835</td>
<td>(Pham et al., 2007)</td>
</tr>
<tr>
<td>pTWEETY::&lt;sub&gt;mutY&lt;/sub&gt;</td>
<td>pTWEETY integrating vector carrying mutY: kan&lt;sup&gt;R&lt;/sup&gt;</td>
<td>7322</td>
<td>(Hassim et al., 2015)</td>
</tr>
<tr>
<td>pTWEETY::&lt;sub&gt;nth&lt;/sub&gt;</td>
<td>pTWEETY integrating vector carrying nth: Kan&lt;sup&gt;R&lt;/sup&gt;</td>
<td>6990</td>
<td>(Moolla et al., 2014)</td>
</tr>
<tr>
<td>pSE100</td>
<td>E. coli-Mycobacterium shuttle vector carrying Pmyc1tetO; Hyg&lt;sup&gt;R&lt;/sup&gt;</td>
<td>5538</td>
<td>Ehrt et al. 2005</td>
</tr>
</tbody>
</table>

amp<sup>R</sup>—ampicillin resistance; oriE—origin of replication in E.coli; kan<sup>R</sup>—Kanamycin resistance; hyg<sup>R</sup>—hygromycin resistance.
2.2. **Bacterial Transformation Conditions**

2.2.1 **Preparation of chemically competent *E.coli* (DH5α) cells**

*E. coli* cells were inoculated in 50 ml of LB media and incubated at 37°C with shaking until the culture reached an optical density at a wavelength of 600nm (OD$_{600nm}$) of 0.5. The freshly grown cells were incubated on ice for 15 minutes (min) and centrifuged at 4500 revolutions per minute (rpm) for 5 min at 4°C. The cell pellets were re-suspended in 30 ml TfbI solution (30 mM potassium acetate, 100 mM rubidium chloride, 10 mM calcium chloride, 50 mM manganese chloride, and 15 % v/v glycerol - pH 5.8) and chilled on ice for 15 min. The cells were then harvested by centrifugation at 4000 rpm at 4°C. The pelleted cells were re-suspended in 10 ml TfbII solution (10 mM MOPS, 75 mM calcium chloride, 10 mM rubidium chloride and 15 % v/v glycerol ), dispensed into 1 ml aliquots and stored at -80°C until required.

2.2.2 **Transformation of chemically competent *Escherichia coli* DH5α cells**

One hundred microlitres of chemically competent cells were chilled on ice for 15 min and mixed with 1-2 μg of plasmid DNA. The tubes were incubated in a heating block at 42°C for 90 seconds (heat shock) and then immediately transferred to ice for 2 min. Subsequently, 700 μl of 2XTY (2% tryptone, 1% NaCl, 1% yeast extract) was added to the cell mixture and incubated at 37°C for 1 hour. Transformed cells were spread on LA agar plates containing the appropriate antibiotic and incubated at 37°C overnight to allow for growth of colonies. As a positive control to assess the transformation efficiency, chemically competent cells were transformed with 1 μg of a replicating plasmid pGEM3Zf(+) and 100 μl of a 10-fold dilution series was spread onto LA plates containing 100 μg/ml ampicillin. As a negative control competent cells were spread on LA plates containing ampicillin to ensure that the cells were
not contaminated. The plates were incubated at 37°C overnight or alternately at 30°C for two days when transforming large plasmids.

### 2.3 Electroporation of \textit{M. smegmatis}

\textit{M. smegmatis} cells were grown in 100 ml of 7H9 with shaking at 37°C to an OD\textsubscript{600nm} of 0.6 - 0.8. The cells were harvested at 4000 rpm for 15 min at 4°C and washed three times by re-suspending the pellet in 20 ml of ice cold 10% glycerol. Washed cells were finally re-suspended in 900 µl of cold 10% glycerol and dispensed into three 300 µl aliquots in Eppendorf tubes. One-two µg of plasmid DNA was mixed with 300 µl of electro-competent \textit{M. smegmatis} cells and then transferred to a cooled electroporation cuvette. The competent cells were exposed to an electrical pulse using the GenePulser\textsuperscript{TM} electroporator (Bio-Rad) with the following parameters: 2.5 kV, 1000Ω and 25 µF. Subsequently, 800 µl of 7H9 was added to the cells and incubated for a minimum of 6 hours to overnight at 37°C to allow for recovery and replication of the transformants. The cells were then spread onto 7H10 plates supplemented with appropriate antibiotic/s and incubated at 37°C for 7-10 days. To determine the transformation efficiency, the replicating plasmid pSE100 (Table 2.2) was electroporated into an aliquot of competent cells, a 10-fold dilution series performed and plated onto7H10 plates containing 50 µg/ml of hygromycin. For the negative control, to ensure that the cells were not contaminated, no DNA was added to the cells and processed as described above.
2.4 Molecular biology techniques

2.4.1 Mini-prep plasmid DNA extraction

The mini-prep plasmid DNA extraction method was carried out using protocol described by Maniatis et al., 1982. *E. coli* clones carrying plasmid DNA were grown overnight at 37°C in 1 ml LB with shaking (250 rpm). One ml of the bacterial cells were harvested by centrifugation at 1300 rpm for 15 min and the pellets were re-suspended in 100 µl of solution I (0.5 M glucose, 0.05 M EDTA, 0.1 M NaOH) followed by the addition of 200 µl of solution II (10 M NaOH, 10% SDS). After gentle inversion, 150 µl of solution III (5 M potassium acetate, 11.5% glacial acetic acid) was added and the tubes vigorously mixed. As a white precipitate formed, the tubes were incubated on ice for 10 min and centrifuged at 13000 rpm at room temperature for 10 minutes. The clear supernatant containing the plasmid DNA was precipitated with 500 µl of isopropanol by centrifugation at 13000 rpm for 10 min. The DNA pellet was washed with 70% ethanol and dried for 20 min in a SpeedVac concentrator (5301Eppendorf). The dried DNA pellet was re-suspended in 150 µl sterile distilled water (sdH₂O) to which 1µl of RNase (10 mg/ml) was added and incubated at 42°C for a further 15 min.

2.4.2 Maxi-prep plasmid DNA extraction

*E. coli* bulk plasmid DNA extraction was performed using the Nucleobond kit (Macherey Nagel) according to the manufacturer’s specifications. *E. coli* clones were grown in 50 ml of LB broth with the appropriate antibiotics and the cultures were harvested by centrifugation at 4000 rpm for 10 min. The cells were re-suspended in 4 ml of buffer S1. Thereafter, 4 ml of buffer S2 was added followed by gentle inversion 5 times. Buffer S3 was added to the mixture, inverted till a white flocculent was visible and the suspension was centrifuged for 25
min at 4500 rpm. A Nucleobond column was equilibrated with 2.5 ml of buffer N2 and the clear supernatant was passed through the column. Thereafter, 10 ml of buffer N3 was added to wash the column and finally the DNA was eluted with 5 ml of buffer N5. The eluent was precipitated with 3.5 ml of isopropanol by centrifugation at 13000 rpm for 30 min. The pelleted DNA was washed with ice cold 70 % ethanol, air dried and dissolved in 200 µl of nuclease free water.

2.4.3 Small scale genomic DNA extraction

For small scale extraction of genomic DNA from *M. smegmatis*, the colony boil method was used. Briefly, a single colony was picked from solid media (7H10) and suspended in 50 µl of sdH2O to which 50 µl of chloroform was added and boiled at 65°C for 15 min. Thereafter, the mixture was centrifuged at 13000 rpm for 15 min and the supernatant containing the DNA was transferred into a fresh Eppendorf tube and used as DNA template for PCR reactions (Section 2.4.6).

2.4.4 Large scale genomic DNA extraction

The cetyltrimethylammonium bromide (CTAB) method (Doyle, 1987) was used for genomic DNA extraction from *M.* strains. Bacterial strains were spread onto solid media (7H10) and incubated at 37°C for two days. Cells were scraped off the plate using an inoculating loop and re-suspended in 500 µl of TE buffer (10 mM Tris-HCl, 10 mM EDTA, pH 8.0) and incubated at 65°C for 35 min. A 50 µl aliquot of lysozyme (10 mg/ml) was added and the mixture incubated at 37°C for 60 min, followed by the addition of 6 µl of proteinase K (10 mg/ml) and 70 µl of 10% SDS. This mixture was incubated at 65°C for a further 120 min. Following the incubation, 100 µl of 5M NaCl and 80 µl of pre-warmed 10% CTAB was
added, mixed gently by inversion, and incubated at 65°C for 10 min. Thereafter, 750 μl of chloroform: isoamyl alcohol (24:1) was added, gently inverted and centrifuged at 13000 rpm for 5 min. The top aqueous phase was transferred to a new tube and 600 μl of isopropanol was added. The DNA was concentrated by centrifugation for 25 min at 13000 rpm and the pellet was washed with 70% ethanol, air dried and dissolved in 500 μl of sdH2O.

2.4.5 Quantifying DNA

The Nanodrop ND-100 Spectrophotometer (NanoDrop technologies) was used to measure the concentration of nucleic acids at an absorbance of 260 nm (A$_{260nm}$). Water was used to blank the machine and 1μl of the DNA suspension was spotted onto the pedestal to measure the quantity and quality (A260/A at 280) of DNA.

2.4.6 Polymerase chain reaction (PCR)

PCR reactions were carried out using the FaststartTaq DNA Polymerase kit, (Roche Applied Science) following the manufacturer’s instructions. All primers used in this study are listed in Appendix 5.2 and Table 5.1 (Moolla et al., 2014; Hassim et al., 2015). All PCR reactions were made up to a final volume of 50 μl, which contained 1-5 μl of template DNA, 2.5 μl each of the 10 mM forward and reverse primers, 10 μl of 10X GC rich buffer, 5 μl of Taq Buffer with 2 mM MgCl$_2$, 0.5 μl HotStartTaq and 8 μl 5XdNTPs(0.2 mM). Sterile distilled nuclease free water was used to adjust the reaction volume. Cycling conditions were carried out with the following parameters: denaturation at 95°C at 5 min, followed by 30 cycles of denaturation at 94°C for 30 seconds (sec), an annealing step at 61-67°C (dependent on primers) for 30 sec and extension at 72°C for 30 sec, ending with a final extension at 72°C for 10 min.
2.4.7 Restriction endonuclease digestions

Restriction endonucleases and their respective buffers were purchased either from New England Biolabs, Roche Biochemicals or Fermentas and used according to the recommendations of the supplier. A standard restriction endonuclease digestion was carried out in a 20 μl reaction containing 0.5-1 μg of DNA and 1X restriction buffer. Bovine serum albumin (BSA) was added where necessary and an appropriate amount of sterile distilled water added to adjust the reaction volume. For plasmid digestions, the reactions were incubated at 37°C for 60 min (or for 30 min if the enzymes displayed star activity) and overnight for genomic DNA.

2.4.8 Agarose gel electrophoresis

DNA fragments were separated on 0.8% - 1% agarose gels. Generally, 5 μl of the DNA sample was mixed with 5μl of loading dye (1 ml 0.5 M Tris-HCl, 0.8 ml glycerol, 1.6 ml 10% SDS, 0.4 ml β-mercaptoethanol, 0.4 ml bromophenol blue and loaded into the wells of the agarose gel with an appropriate molecular weight marker (Roche Applied Science) for approximate size determination. The DNA fragments were separated in 1X TAE buffer (1 mM EDTA, 40 mM Tris-acetic acid) at 90 V for 30 to 45 min depending on the size of the DNA fragments. For Southern blot analysis, the overnight genomic DNA digestion was separated for 2 to 3 hours on a 0.7% agarose gel at 80 V in cold TAE.

2.4.9 DIG labelled probe synthesis

To generate a labelled probe for Southern blot analysis, a PCR DIG Probe synthesis kit (Roche Applied Science) was used as per the manufacturer’s instructions. Briefly,
approximately 1 μg of appropriate plasmid DNA was diluted to 50 ng with dH₂O. The PCR was carried out in a total volume of 50 μl with 1 mM forward and reverse primers added as listed in Table 5.1(Appendix 5.2) to amplify the upstream and downstream homologous regions. The positive PCR reaction was supplemented with 5 μl of DIG labelling mix and no DIG label was added to the negative control. The PCR product sizes were analysed on a 1% agarose gel to determine incorporation of the DIG label; the labelled product runs at a higher molecular weight on the gel compared to the unlabelled control, since incorporation of DIG increases the molecular size of the product. Five μl of the synthesized probe was added to 15 ml of DIG Easy Hyb solution (Roche Applied Science) for Southern blotting (Section 2.4.10) and the remaining labelled probe was stored at -20°C.

2.4.10 Southern blot hybridizations

The Southern blot technique enables a specific DNA sequence of interest to be identified from a complex mixture of DNA (Southern, 1975). This technique involves the separation of digested DNA by gel electrophoresis, transfer to a nitrocellulose membrane and labelling of the target DNA using a DIG labelled probe for visualization. Briefly, genomic DNA was prepared using the CTAB method (Section 2.4.4) from culture plate of M. smegmatis (mc²155, Δnth, ΔmutY, SCO strains, ΔmutYΔnth and ΔnthΔmutY Table 2.1). Approximately 2–3 μg of the genomic DNA was digested with appropriate enzymes and incubated overnight at 37°C. The digested DNA was separated on a 0.7% agarose gel as described in Section 2.4.8. The agarose gel was soaked in depurination buffer (0.2 M HCl) for 15 min at room temperature with gentle shaking, followed by two rinses with distilled water. The gel was then immersed in denaturation buffer (0.5 M NaOH, 1.5 NaCl) for 30 minutes, after which it was washed twice in distilled water and equilibrated in 1X TBE solution (89 mM Tris-Borate, 2 mM MEDTA, pH 8.3). The agarose gel was then covered with a Hybond
nitrocellulose membrane (Amersham) that was cut to size and sandwiched between two pieces of filter paper and two sponges. The assembled layers were enclosed in a plastic cassette and transferred to an electroblotting unit (OmniPAGE Electroblotting Unit, Cleaver Scientific Ltd CS-300 V) containing 1X TAE. The DNA was transferred onto the nitrocellulose membrane at 10 V for 2 hours at 4°C. Subsequently, the DNA was UV cross-linked onto the nitrocellulose membrane at 2500 mJ/cm² in a UV-crosslinker for 2 minutes (UV Stratalinker 1800, Stratagene).

The membranes were pre-hybridized with 10 ml of DIG Easy-Hyb at 55°C for 15 min in a hybridizing oven (Hybrid Micro-4). The DIG Easy-Hyb solution with DIG labelled probe was heated to 95°C for 5 min and added to the hybridization bottle with the membranes. The membranes were hybridized overnight at 55°C in hybridizing oven. The following morning, the membranes were washed in 20 ml of pre-warmed 2X SSC wash solution I (300 mM NaCl, 30 mM sodium citrate and 0.1 % SDS), for 5 min at room temperature. Thereafter, the membranes were washed in solution II (0.5 × SSC and 0.1 % SDS) in the hybridizing oven at 65°C for 15 min. The above process was repeated twice after which the membranes were rinsed with wash buffer (0.1 M maleic acid buffer and 0.3 % Tween 20) for 10 min with gentle agitation at room temperature. Two microlitres of Anti-Digoxigenin-AP (Roche Applied Science) supplied with the kit was added to 20 ml of blocking solution and the membranes incubated for 30 min with gentle agitation. The membranes were then incubated in detection buffer for 5 min and placed inside a plastic hybridization bag (Roche Diagnostics; Mannheim, Germany) to which 1 ml CSPD (which emits a luminescent signal) was added. The hybridization bag was heat sealed and incubated for 10 min at 37°C. The membrane was then placed in an imaging cassette, exposed to X-ray film and placed in a dark
drawer for approximately 30-60 min. The X-ray film was developed using an automated developer (Axim).

2.5 Knockout vector construction

Plasmid constructs containing inactivated alleles of the nth and mutY genes were previously generated by PCR (Moolla, et al., 2014; Hassim, et al., 2015). Briefly, as shown in Figure 4 (1) approximately 1500 bp upstream and downstream (red and green blocks respectively) of the genes of interest, containing residual 3’ and 5’ ends of the gene, were amplified by PCR using a high fidelity polymerase and primers that had appropriate restriction sites incorporated within the sequence. (2). The amplified upstream and downstream fragments were cloned into pGEM3Zf(+) (Appendix 5.3, Figure. 30) and sequenced to ensure mutations were not introduced during the PCR cycling. Subsequently, the upstream and downstream fragments were excised from pGEM3Zf(+) and cloned by three way cloning into the suicide vector, p2NIL (Appendix 5.3, Figure 31) which carries a kanamycin resistance marker. The Pac cassette from pGOAL17 (Appendix 5.3 Figure, 32) containing marker genes sacB and lacZ was inserted into the PacI site of p2NIL (Figure 4 [4]) to generate p2NILΔmutY::pGOAL17 (map shown in Figure 10) or the Pac cassette from pGOAL19 (Appendix 5.3 Figure. 33) with sacB, lacZ, hyg and was ligated into the PacI site in p2NIL to generate p2NILΔnth::pGOAL19 (map shown in Figure 9).
Figure 4. Schematic representation of the construction of knockout constructs using three way cloning. US – upstream region (red), DS – downstream region (green). The bla gene confers ampicillin resistance, aph gene confers kanamycin resistance and hyg gene confers hygromycin resistance. The black arrows represent forward and reverse primers.
2.5.1 Inactivation of the Nth and MutY DNA glycosylases to generate double deletion mutants

Homologous recombination using two-step allelic exchange methodology as described by (Gordhan and Parish, 2001) was used to generate gene deletion mutants. This involved the use of a suicide vector p2NIL (Table 2.2) carrying a truncated version of either the nth or mutY gene (Table 2.1) which was electroporated into competent M. smegmatis cells. Through homologous recombination the non-functional gene and the vector backbone is integrated into the mycobacterial genome, generating a single crossover mutant (SCO) carrying the wildtype gene, truncated gene and marker genes (aph, sacB and lacZ). The lacZ gene, which expresses the β-galactosidase enzyme that degrades X-gal into a blue colour, results in blue colonies which allows for the selection of single cross-over (SCO) homologous recombinants which carrying a copy of the functional gene and a copy of the truncated gene as depicted in Figure 5 (1 and 2). Single cross-overs were allowed to undergo a double cross over event in the presence of sucrose. The basis for this is that colonies retaining the sacB cassette will not be viable in the presence of sucrose, since sacB expresses levansucrase which converts sucrose to toxic levan thus killing the bacteria. Hence, white sucrose resistant colonies lose all the marker genes together with the vector sequence and are indicative of successful double cross over (DCO) transformants as shown in Figure 5 (3).
2.5.2 Generation of SCO and DCO transformants

The p2NILΔmutY::pGOAL17 and p2NILΔnth::pGOAL19 suicide plasmids were electroporated into mid-logarithmic phase cultures of the Δnth and ΔmutY deletion mutant strains respectively and selected on 7H10 plates containing kan and X-gal to identify for SCO’s. The genotype of the SCO’s was confirmed by PCR (Section 2.4.6) and Southern blot analysis (Section 2.4.10). For the PCR, chromosomal DNA was extracted from potential SCO’s using the colony boil method (Section 2.4.3) and the DNA amplified using primers that distinguished the mutant and the wild type alleles (Appendix 5.2 Table 5.1).

To generate DCO mutants, one of the PCR confirmed SCO’s was selected and grown overnight in 10 ml 7H9 with kan. The overnight culture was inoculated into 20 ml of fresh 7H9 without antibiotics and grown at 37°C overnight to allow for the second crossover to

Figure 5. Diagrammatic representation of allelic replacement by homologous recombination. Marker genes – sacB, lacZ, aph and/or hygromycin resistance. Modified from (Gordhan and Parish, 2001).
take place. The cells were serially diluted (10-fold) and 100 μl of each of the dilutions (10^0 - 10^8) were plated on 7H10 supplemented with 2% sucrose and X-gal. Plates were incubated at 37°C and scored for white sucrose resistant colonies, which were potential DCO mutants. Potential DCO’s were further confirmed for kan sensitivity by spotting on 7H10 media with and without the antibiotic. The potential confirmed DCO’s were screened initially by PCR to distinguish between wild type and mutant alleles (Figures 11 and 12). The PCR confirmed putative mutants together with the SCO and parental stain were further analysed by Southern blot to ensure that the genes upstream and downstream of the deletion were genotypically intact.

### 2.5.3 Complementation of M. smegmatis mutants

To confirm that the phenotypes observed for the double gene knockout mutant strains was due to the loss of the respective DNA glycosylase gene and not due to downstream polar mutations, complementation strains were generated. For complementation, the full length of each functional gene with 350 bp of upstream sequence, carrying the promoter region, was previously amplified and cloned into the integrating vector pTWEETY (Appendix 5.3 Figure 33) to generate pTWEETY::nth and pTWEETY::mutY (Table 2.1). Both these constructs were created by two previous Masters students (Moolla et al., 2014, Hassim et al., 2015). As a secondary confirmation these vectors were rescreened in this study by restriction analysis to confirm that the vectors did not rearrange. The appropriate complementation vector was electroporated into the respective mutant strain and the resulting clones were screened by PCR (Figure 19) using the colony boil method for DNA extraction. Two complemented strains, ΔmutYΔnth::nth and ΔnthΔmutY::mutY were generated (Table 2.1).
2.6 Phenotypic characterization of DNA glycosylase deficient mutant strains

2.6.1 Growth kinetics under normal culture conditions
To determine growth kinetics of the various mutant strains, 1 ml bacterial freezer stocks were grown overnight at 37 ºC in 5 ml 7H9 medium with shaking at 100 rpm. The overnight late log phase culture was diluted into 50 ml of fresh media to a starting $\text{OD}_{600\text{nm}}$ of 0.02 and growth of the cultures was monitored by $\text{OD}_{600\text{nm}}$ measurements every 3 hours for a period of 36 hours. Growth was represented graphically as optical density over time.

2.6.2 Mutation frequency
For mutation frequency analysis, bacterial strains were grown overnight in 20 ml 7H9 at 37 ºC to an $\text{OD}_{600\text{nm}}$ of 0.5 - 0.8. Aliquots (100µl) of 10 fold serial dilutions ($10^4$-$10^7$) were spread onto 7H10 media for colony forming unit (CFU) counts. The remaining culture was centrifuged for 15 minutes at 4500 rpm and the pellet re-suspended in 500 µl of fresh 7H9 broth. The cells were spread onto 7H10 supplemented with 200 µg/ml rifampicin (rif) and incubated at 37ºC. The plates were scored after 7 days for rifampicin resistant (rif$^R$) colonies.

Mutation frequency was calculated as total number of rif$^R$ colonies divided by the total number of colony forming units (CFU/ml) per culture.

2.6.3 UV induced mutagenesis
Overnight cultures grown in 100 ml of 7H9 to an $\text{OD}_{600\text{nm}}$ of 0.6 in a shaking incubator were harvested by centrifuging at 4000 rpm for 15 minutes. The cells were re-suspended in 12 ml of 7H9 and split into two aliquots of 6 ml each. One aliquot was treated with UV at 50 mJ/cm$^2$ (UV Stratalinker 1800, Stratagene) and the other aliquot was the untreated control. 1
ml of the untreated cells was plated onto 7H10 plates containing 200 µg/ml rif and 0.1 ml for each of the dilutions (10^5-10^8) was plated onto 7H10 for viable colony counts. After UV treatment, damaged cell were added to 50 ml 7H9 and allowed to recover for 6 hours. At 3 and 6 hour time intervals, a 2 ml aliquot was sampled, serially diluted (10^5-10^8) and spread onto 7H10. The plates were incubated at 37°C for 3 days before scoring. In addition 1ml was spread onto 7H10 rif (200 µg/ml) plates and scored after 7 days for rif^R colonies. The induced damaged mutation frequency was calculated as total number of rif^R colonies divided by total population of cells (CFU/ml). The survival of UV treated cultures was scored by counting CFU of the surviving cells and represented graphically as log CFU/ml over time.

2.6.4 Sensitivity to H_2O_2

Bacteria were grown in 30 ml of 7H9 at 37°C to early log phase (OD_{600nm} 0.35) and hydrogen peroxide (2.0 mM) was added to the cells to induce oxidative stress. At 2, 4 and 6 hour time intervals, 1 ml aliquots were removed, serially diluted (10^0 -10^7) in 7H9 and 0.1 ml of each dilution spread onto 7H10 plates. The plates were incubated at 37°C for 3 days before scoring for CFU’s of surviving cells. The surviving cells were represented graphically as log CFU/ml over time.

2.6.5 H_2O_2 diffusion susceptibility test

To investigate the sensitivity of the wildtype, single and double mutants and their respective complemented strains to H_2O_2, cultures of each of the strains were grown overnight in 7H9 at 37°C with shaking to an OD_{600nm} of 0.35. A hundred microliters of the overnight culture was plated on 7H10 and allowed to dry for an hour. Aliquots of 0.5, 1.0 and 1.25 µl of an 880 mM
stock solution of H$_2$O$_2$ (0.176, 0.352 and 0.440 mM final concentration in agar plates respectively) was spotted in the centre of the plate and incubated at 37°C for three days. Zones of inhibition were recorded as the distance between the cleared zone and the edge where there was visible bacterial growth.

2.6.6 Mutation rate measurements

A mutation rate measure the likelihood of a cell gaining a mutation during its lifetime and takes into account the possibility of mutations arising either early or late during growth. The Luria-Delbrück fluctuation analysis was used to measure spontaneous mutation rates for the various mutants. This method allows for the number of mutational events to be accurately calculated by using parallel cultures, thereby minimizing variability (Rosche and Foster, 2000). The mutation rate was calculated as described previously by Machowski et al., 2007. Briefly, mutation rate is taken as the number of observed mutations per culture ($m$) within the total population ($Nt$). The mutation rates were analysed using either the P naught method (P0) or the Lea-Coulson Maximum Likelihood method (MSS-LC) (Rosche and Foster, 2000). To decide on the method of choice, the number of rif$^R$ mutants per culture ($m$-value) needed to be determined. In order to estimate the $m$-value, the Ma-Sandri-Sarkar Maximum Likelihood Estimator (MSS-MLE) and the $Po$ method were combined as explained by the Luria-Delbrück distribution. Using a spreadsheet for calculations generated by Machowski et al., 2007, MSS-LC distribution graph/curve was generated (Figure 6). The peak (green arrow) of the curve is the determined as estimated $m$-value which used to calculate the mutation rate. The P0 method was used when there was a low number of rif$^R$ mutants on selective plates, in this case it was used for analysis of the wildtype, Δnth, ΔmutY and their respective complemented strains as the $m$-value was between 0.3 and 2.3. The MSS-LC was used when a high number of rif$^R$ mutants ($m$-value) were obtained on selective plates, which was used
for ΔnthΔmutY, ΔmutYΔnth and their respective complemented strains as them-values were between 1 to 15. The Mutation rate was calculated using the following formula: 

\[ \mu = \frac{m \times 10^2}{Nt-value} \]

Where \( m \) is the number of observed mutations per culture and \( Nt \) the total final number of cells in a culture. Frozen stocks of the parental, mutant and complemented strains were streaked on 7H10 and incubated at 37°C until single colonies emerged. Single colonies from the respective strains were grown in 7H9 media to late log phase (10^8 cells/ml). A dilution series of the culture was performed and spread on 7H10 media to measure the initial inoculum (\( N_0 \)). Also, 1 ml of the culture was spread onto 7H10 rifampicin plates to ensure that there were no pre-existing mutants. The cell density was diluted from 10^8 cells/ml to 10^2 cells/ml and 2 ml aliquots were dispensed into 30 culture tubes which were incubated at 37°C with constant shaking (100 rpm) for 6 to 7 days until the cultures were in late log phase. The total population size (CFU/ml) was determined by randomly selecting 5 culture tubes out of the 30 and spreading a serial dilution of these on 7H10 plates (\( Nt \)). The culture from the remaining 25 tubes was plated individually on 7H10 rif (200 µg/ml) to score for spontaneous rif^R mutants (\( m \)-value). The 7H10 plates were incubated at 37°C for 3 days for total population size count and 7 days for scoring rif^R mutants.
Figure 6. An example of the MSS-LC distribution graph/curve use to estimate the Rif$^R$ mutants per culture ($m$-value). The green arrow represents the peak of the graph which is the estimated $m$-value from the data of the experiment.

### 2.6.7 Spectral analysis of rifampicin resistant mutants in the *rpoB* region

Most of the spontaneous rifampicin resistant mutations in mycobacteria are associated with mutations within the 81 bp rifampicin resistance-determining region (RRDR) found in the *rpoB* gene (Siu *et al*., 2011). To assess the spectrum of mutations in the rifampicin resistance-determining region (RRDR) of the *rpoB* gene, genomic DNA was extracted using the colony boil method from rifampicin resistant colonies isolated from the fluctuation assay (Section 2.6.6). Primers MSmrBR2 and MSmrBF2 listed in. (Appendix 5.2 Table 5.1) were used to amplify the RRDR of *rpoB* (549 bp). The PCR products were sequenced by DNA Sequencer Central Analytical Facility in Stellenbosch using the sequencing primer MSmrBF1 (Appendix 5.2 Table 5.1) to identify mutations resulting in rif$^R$. 

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![Figure 6. An example of the MSS-LC distribution graph/curve use to estimate the Rif$^R$ mutants per culture ($m$-value). The green arrow represents the peak of the graph which is the estimated $m$-value from the data of the experiment.](image-url)
3. Results

3.1 Bioinformatics of Nth and MutY DNA glycosylases

Bacterial survival in an oxidative environment requires constant adaptation. Bacteria are equipped with proteins and genes that detoxify, scavenge and repair the DNA damage caused by endogenous oxidative stress. When there is damage, glycosylases in the base excision repair (BER) pathway are responsible for locating and removing oxidatively damaged bases. Studies in *E. coli* of the Nth and MutY glycosylases clearly identified a functionally unique role for each enzyme however, some level of structural similarity has been recognized between the two proteins (Thayer *et al*., 1995). Crystal structures of the *E. coli* Nth and MutY DNA glycosylases show that both these proteins have the Helix-hairpin-Helix domain which serve as a DNA binding motif and a highly conserved 4Fe-4S domain, also important for DNA binding (Guan *et al*., 1998, Kuo *et al*., 1992). Previous mycobacterial amino acid alignments of Nth and MutY glycosylases identified that they contain an iron-sulfur cluster with the four cysteine residues and Helix-hairpin-Helix domain (Moolla, *et al*., 2014; Hassim, *et al*., 2015). In *M. tuberculosis*, *M. smegmatis* and *E. coli*, the nth gene is in the same chromosomal context (Moolla, *et al*., 2014). Comparisons of the protein crystal structures of the *E. coli* Nth and MutY revealed a 66% similarity between the two enzymes (Michaels *et al*., 1990). Similarly using tertiary structure prediction software (http://zhanglab.ccmb.med.umich.edu/I-TASSER/), the protein structures of both Nth and MutY from *M. smegmatis* were modelled (Figures 7A and 7B). The predicted protein structures showed the presence of the Helix-hairpin-Helix (HhH) motif, the four iron-sulphur (4Fe-4S) cluster for both the glycosylases, which appear to be highly conserved in mycobacteria. The overall folding of the predicted protein structures for both the enzymes are similar, with most of the residues buried inside the globule like structure (Figures 7A and 7B). Interestingly, both proteins in mycobacteria have conserved the helix-hairpin-helix and
the 4Fe-4S DNA binding residues. The one difference between the two enzymes is that at position 120, the tyrosine (Tyr120) in MutY (Figure 7A) is replaced with lysine (Lys) in the Nth DNA glycosylase (Figure 7B), suggesting that the latter enzyme has lyase activity whilst MutY does not (shown in turquoise).

Figure 7. Predicted protein structure of the M. smegmatis Nth and MutY DNA glycosylases. Proteins were modelled using the I-TASSER pymol software (http://zhanglab.ccmb.med.umich.edu/I-TASSER/). (A) MutY DNA glycosylase and (B) Nth DNA glycosylase.

Amino acid comparisons of the sequences of the Nth and MutY proteins from M. tuberculosis, E. coli and M. smegmatis was done to confirm the amino acids responsible for lyase activity and to identify residues which may play an important role in determining substrate specificity (Figure 8). A pairwise alignment of the Nth and MutY glycosylases of M. smegmatis demonstrated 17% identical with 28% similar on the primary amino acid level, and for M. tuberculosis the proteins are 30% similar with 20% identical. In E. coli the Asp 138 residue in both Nth and MutY has been shown to be important for catalytic activity (Yang et al., 2000). The Lys120 residue, has been reported to be essential for AP lyase
activity and cleaves the glycosidic bond by nucleophilic attack, noticeably this residue is replaced with valine in the *E. coli* MutY hence, accounting for the lack of mechanistic lyase activity (Thayer *et al.*, 1995, Guan *et al.*, 1998). The amino acid alignments (Figure 8) show that in both *M. smegmatis* and *M. tuberculosis* the Nth DNA glycosylase has retained the lysine residue at position 120, but in the MutY protein it has been replaced with tyrosine (Tyr120), suggesting that MutY in mycobacteria acts only as a glycosylase and similar to the *E.coli* counterpart does not have lyase activity.
**Figure 8.** Amino acid sequence alignments of the Nth and MutY DNA glycosylases from *M. smegmatis*, *M. tuberculosis* and *E. coli*. The helix is shaded dark yellow, the hairpin is shown in red, the aspartic acid residue is in blue (position 148), the arginine residue is shown in pink at position 153, which is conserved in both proteins, the cysteine residues involved in the binding of the 4Fe-4S cluster is shaded in green. The conserved lysine residue in Nth (but not in MutY) at position 120, responsible for lyase activity, is shaded in turquoise.
3.2 Confirmation of existing knockout constructs

In order to ensure that the deletion constructs for *nth* and *mutY* generated in previous studies (Moola *et al* 2014; Hassim *et al* 2015) were correct, DNA from the *E. coli* strains carrying the p2NIL*mutY::pGOAL17* and p2NIL*nth::pGOAL19* knockout vectors, were isolated and digested with various restriction enzymes. Both constructs yielded in the corresponding expected sizes, thus confirming that the knockout vectors were correct (Figures 9 and 10).

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Expected sizes(bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Sph</em>I</td>
<td>14742</td>
</tr>
<tr>
<td><em>Hind</em>III</td>
<td>9451, 4525, 739, 27</td>
</tr>
<tr>
<td><em>Kpn</em>I</td>
<td>6803, 5197, 1580, 1132, 30</td>
</tr>
<tr>
<td><em>Pvu</em>II</td>
<td>7498, 3259, 2557, 819, 363, 144, 102</td>
</tr>
<tr>
<td><em>Nru</em>I</td>
<td>4954, 4356, 3470, 1061, 412, 270, 219</td>
</tr>
</tbody>
</table>

**Figure 9.** The p2NIL*nth::pGOAL19* deletion construct. (A) Table displaying the different restriction endonucleases used and the expected fragment sizes (bp), (B) Plasmid map of the p2NIL*nth::pGOAL19* vector showing the cloned US and DS regions (shaded in green and blue respectively) and (C) Agarose gel of digested fragments of p2NIL*nth::pGOAL19* vector.
Figure 10. The p2NILmutY::pGOAL17 deletion construct. (A) Table displaying the different restriction endonucleases used and the expected fragment sizes, (B) Plasmid map of the vector showing the cloned US and DS regions (shaded in green and blue respectively) and (C) Agarose gel of digested fragments of the p2NILΔmutY::pGOAL17.

3.3 Generation and identification of single crossover (SCO) and double crossover (DCO) mutants

Suicide vectors p2NILΔnth::pGOAL19 and p2NILΔmutY::pGOAL17 were electroporated into previously generated mutant strains lacking either mutY (ΔmutY) or nth (Δnth) (Moolla, et al., 2014; Hassim, et al., 2015) respectively to generate double gene knockout mutant strains. After electroporation, blue transformants (SCO mutants) selected on 7H10 agar plates.
supplemented with kanamycin (kan) and X-gal were screened for genotypic confirmation initially by PCR followed by Southern blot analysis (Section 3.3.2). To achieve the second crossover, PCR verified SCO mutants were cultured in non-selective media and plated on 7H10 plates containing 2% sucrose. White, sucrose resistant clones resulting from the loss of the marker genes on the suicide vector during the double cross over event selection were screened by PCR to identify positive double cross over (DCO) clones.

3.3.1 PCR confirmation of double mutants

PCR primers used for screening the nth and mutY deletion region were designed previously and are listed in (Appendix 5.2 Table 5.1). The primers were designed to bind upstream and downstream of the gene to distinguish the wildtype allele from the mutant allele hence, a smaller PCR product was amplified for the deleted mutant allele. The expected size for the nth deletion using the primer set NthF1and NthR1 (Appendix 5.2 Table 5.1) was 270 bp and 833 bp for the wildtype allele. For the mutY deletion a 780 bp fragment was expected for the deletion allele and a 1481 bp fragment for the parental allele using the primer set mutYFwd and mutYRev (Appendix 5.2 Table 5.1). The PCR data shown in Figure11 and 12 identified several possible nth (6) and mutY (7) deletion double mutants in the ΔmutY and Δnth background strains respectively.
Figure 11. PCR screen for identification of double gene knockout deleted mutant strains using nth primers. (A) Schematic diagram showing screening primers (red arrows) with forward primer (F1) located upstream of the gene and reverse primer (R1) located downstream of the gene. (B) Agarose gel showing the PCR screen for deletion of nth in the ΔmutY deletion mutant. [Lane 1] Marker VI, [Lane 2] mc^2 155, [Lane 3] p2NILΔnth::pGOAL19 plasmid DNA, [Lane 4] SCO mutant strain, [Lanes 5 to 14], possible DCO ΔmutYΔnth clones. Expected sizes, for wild type (mc^2 155) =833 bp and for Δnth =270 bp.
Figure 12. PCR screen for identification of double gene knockout deleted strain using mutY primers. (A) Schematic diagram showing screening primers (red) with forward primer (F1) located upstream of the gene and reverse primer (R1) located downstream of the gene. (B) Agarose gel showing the PCR screen for deletion of mutY in the Δnth deletion mutant. [Lane 1] Marker VI, [Lane 2] mc^2155, [Lane 3] SCO strain, [Lane 4] p2NILΔmutY:pGOAL17, [Lane 5 to 15], possible DCO ΔnthΔmutY clones. Expected sizes, for wild type (mc^2155) = 1481 bp and for the ΔmutY =780 bp.

3.3.2 Southern Blot confirmation of double mutants

The PCR confirmed potential mutants where further screened by Southern blot analysis to validate the integrity of the upstream and downstream regions flanking the deletion to ensure that the possible deletion mutants were genetically intact and no chromosomal rearrangement occurred, in addition to confirming integration at the correct site within the genome. To confirm the presence of nth in the wildtype and complemented strains and its deletion in the
respective mutant backgrounds; genomic DNA from the parental strain mc\textsuperscript{2}155, the single \(\Delta\text{mutY}\) on \(\Delta\text{nth}\) mutant strains, the SCO’s obtained during the generation of the double deletion mutants and the possible double deletion mutant strains (\(\Delta\text{mutY}\Delta\text{nth}\) and \(\Delta\text{nth}\Delta\text{mutY}\)) was extracted. For genotypic confirmation of the \(\Delta\text{mutY}\Delta\text{nth}\) mutants, the genomic DNA was digested with \textit{PstI} for analysis of the region upstream of the deletion and with \textit{BamHI} to check the downstream region (Figures 13 and 14). Similarly, the presence of \textit{mutY} in the wildtype and complemented strains and its deletion in the mutant strains was assessed by digesting the genomic DNA with \textit{EcoRI} for the upstream region and \textit{BglIII} for the downstream region from the same strains listed above (Figures 15 and 16).
Figure 13. Genotypic characterization of mc²155, ΔmutY, SCO and ΔmutYΔnth by Southern blot analysis. The genomic map of the relevant locus is shown for mc²155, ΔmutY, SCO and ΔnthΔmutY. Also shown below is the Southern blot. Chromosomal DNA from mc²155, ΔmutY, SCO and ΔmutYΔnth was digested with PstI (P). The probe used for hybridization (upstream probe [US]) is shown by the solid black box and the expected sizes are indicated with a black arrow. The nth gene is highlighted in purple. Note that in mc²155 and ΔmutY the genomic content around Δnth is the same. The single deletion mutant, ΔmutY was previously genotyped for genomic integrity (Hassim et al., 2015) hence, this region is not shown on the maps.
Figure 14. Genotype characterization of mc²155, ΔmutY, SCO and ΔmutYΔnth by Southern blot analysis. The genomic map of the relevant locus is shown for mc²155, ΔmutY, SCO and ΔnthΔmutY. Also shown below is the Southern blot. Chromosomal DNA from mc²155, ΔmutY, SCO and ΔmutYΔnth was digested with BamHI (B). The probe used for hybridization (downstream probe [DS]) is shown by the solid black box and the expected sizes are indicated with a black arrow. The nth gene is highlighted in purple. Note that in mc²155 and ΔmutY the genomic content around Δnth is the same. The single deletion mutant, ΔmutY was previously genotyped for genomic integrity (Hassim et al., 2015) hence, this region is not shown on the maps.
Figure 15. Genotypic characterization of mc²155, Δnth, SCO and ΔnthΔmutY by Southern blot analysis. The genomic map of the relevant locus is shown for mc²155, Δnth, SCO and ΔnthΔmutY. Also shown below is the Southern blot. Chromosomal DNA from mc²155, Δnth, SCO and ΔnthΔmutY was digested with Bg/II (Bg). The probe used for hybridization (upstream probe [US]) is shown by the solid red box and the expected sizes are indicated with a black arrow. The mutY gene is highlighted in green. Note that the genomic content around mc²155 and Δnth is the same. The single deletion mutant, Δnth was previously genotyped for genomic integrity (Moolla et al., 2014) hence, this region is not shown on the maps.
Figure 16. Genotypic characterization of mc^2155, Δnth, SCO and ΔnthΔmutY by Southern blot analysis. The genomic map of the relevant locus is shown for mc^2155, Δnth, SCO and ΔnthΔmutY. Also shown below is the Southern blot. Chromosomal DNA from mc^2155, Δnth, SCO and ΔnthΔmutY was digested with EcoRI (E). The probe used for hybridization (downstream probe (DS)) is shown by the solid red box and the expected sizes are indicated with a black arrow. The mutY gene is highlighted in green. Note that in mc^2155 and Δnth the genomic content around ΔmutY is the same. The single deletion mutant, Δnth was previously genotyped for genomic integrity (Moolla et al., 2014) hence, this region is not shown on the maps.
3.4 Complementation of the double deletion mutant strains

Complementation of ΔnthΔmutY and ΔmutYΔnth with mutY and nth respectively, was accomplished by utilizing previously constructed integrating vectors (Moolla et al., 2014, Hassim et al., 2015). The pTWEETY::nth and pTWEETY::mutY (Table 2.2) are integrating vectors carrying a functional copy of the nth or mutY genes with an additional 350 bp upstream sequence that contains the promoter sequence. To reconfirm genetic integrity of the two integrating vectors, restriction mapping was carried out and the fragments sizes for both pTWEETY::nth and pTWEETY::mutY were confirmed to be correct, Figures 17 and 18. The complementation vector, pTWEETY::nth was electroporated into the double mutant strain ΔmutYΔnth to revert the strain to the ΔmutY single deletion mutant. Similarly, pTWEETY::mutY was integrated into the ΔnthΔmutY mutant to obtain the Δnth single deletion mutant genotype. Transformants were screened by PCR (Figures19 a and b), which confirmed the integration of the respective functional gene into the respective deletion mutants as both the functional and deleted alleles were identified. The resulting complemented strains were annotated as ΔnthΔmutY::mutY and ΔmutYΔnth::nth (Table 2.1). In both the PCR confirmations non-specific bands were observed. This a common observation due to the high G+C rich content in mycobacteria primers often mis-prime resulting in non-specific products,
**Figure 17.** The complemented vector for the *nth* deletion. (A) Table displaying the different restriction endonucleases used and the expected fragment sizes, (B) Plasmid map of the pTWEETY::*nth* vector and (C) Agarose gel of digested fragments of the pTWEETY::*nth* vector.
Table displaying the different restriction endonucleases used and the expected fragment sizes.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Expected sizes (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BglII</td>
<td>4296, 3026</td>
</tr>
<tr>
<td>EcoRI</td>
<td>4466, 2850</td>
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<tr>
<td>HindIII</td>
<td>6355, 961</td>
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<tr>
<td>PstI</td>
<td>7322</td>
</tr>
<tr>
<td>PvuII</td>
<td>2999, 264, 1699</td>
</tr>
</tbody>
</table>

Figure 18. Restriction mapping of the complemented vector for pTWEETY::mutY. (A) Table displaying the different restriction endonucleases used and the expected fragment sizes, (B) Plasmid map of the pTWEETY::mutY vector and (C) Agarose gel of digested fragments of the pTWEETY::mutY vector.
3.4.1 PCR confirmation of complemented strains

Figure 19. PCR conformation of complemented strains. (A) Complementation of ΔnthΔmutY with ΔmutY. [Lane 1] Marker VI, [Lane 2] mc^2155, [Lane 3] Δnth, [Lane 4] ΔnthΔmutY, [Lane 5 to 6] ΔnthΔmutY::mutY. Expected sizes, mc^2155 (1481 bp) and Δnth (1481 bp), ΔnthΔmutY (780 bp) and ΔnthΔmutY::mutY (1481 bp and 780 bp). (B) Complementation of ΔmutY Δnth with nth, [Lane 1] Marker VI, [Lane 2] mc^2155, [Lane 3] ΔmutY, [Lane 4] ΔmutYΔnth, [Lane 5 to 6] ΔmutY Δnth::nth. Expected sizes, wildtype (833 bp), ΔmutY (833), ΔnthΔmutY (270 bp) and ΔmutYΔnth::nth (833 and 270 bp).

3.5 Phenotypic characterisation

3.5.1 Growth kinetics

Growth kinetics of mc^2155, single mutants, double mutants and complemented strains were assessed under normal culture conditions to evaluate the effect of the combinatorial loss of nth and mutY in the ΔmutYΔnth and ΔnthΔmutY strains respectively. Freezer stocks of the
various strains were inoculated in 7H9 media and bacterial growth monitored by measuring the OD$_{600nm}$ at 3 hours intervals over a period of 36 hours. Both the double mutant strains showed comparable growth to the parental strain and the respective single deletion mutants suggesting that Nth and MutY are dispensable for growth in vitro. Furthermore, both the complemented strains also displayed the same growth kinetics as the respective single gene deletion mutants indicating that integration of the functional gene at the attB site does not affect the growth kinetics of these strains.

Figure 20. Growth curve of *M. smegmatis* mc$^2$155, single mutants, double mutants and complemented strains under normal culture conditions. The represented data points are an average of three independent experiments and the standard error for each point is included.
3.5.2 Spontaneous mutation frequency

The parental, mutant and complemented strains were grown overnight to approximately $2 \times 10^7$ CFU’s/ml and 500 µl was plated onto rifampicin (200 µg/ml). The data in Figure 21 and 22 showed that the deletion of mutY in the Δnth background (ΔnthΔmutY) resulted in an exaggerated increase in the level of spontaneous rifR mutations. Comparatively, the Δnth, ΔmutY and ΔmutYΔnth strains showed marginal increase in mutagenesis compared to mc2155. Both the nth complemented strains reduced the mutation frequency to the level of the respective parental strains, whereas the ΔnthΔmutY::mutY strain showed a slightly more mutants compared to Δnth. A possible interpretation of this observation is that the Nth DNA glycosylase plays a greater role in controlling mutagenesis hence, its deletion, prior to deletion of mutY results in exacerbated spontaneous rifR mutants when compared to the ΔmutYΔnth strain, where the order of the deletion of the two glycosylases is reversed. These data suggest a possible hierarchy between the Nth and MutY DNA glycosylases in the maintenance of mycobacterial genome integrity.

**Figure 21.** Spontaneous RifR mutation for single mutants, double mutants and complemented strains.

The error bars represent standard error between 3 biological replicates.
**Figure 22.** Rifampicin containing plates showing rif\(^R\) mutants from spontaneous mutation frequency assay. The plates are representatives of one of 3 experiments. The plates are representatives of one of 3 experiments. The increased rifampicin resistant colonies observed for \(\Delta nth\Delta mutY\) (red box) was consistent over 3 replicates which is markedly different when compared to the \(\Delta mutY\Delta nth\) double mutant strain (green).

### 3.5.3 Effect of H\(_2\)O\(_2\) on the survival of *M. smegmatis*

Mycobacteria are subjected to host-derived ROS such as hydroxyl radical (•OH) and H\(_2\)O\(_2\) which have the ability to cause damage to DNA through oxidative stress. In this study, H\(_2\)O\(_2\) was used as a source of inducing oxidative stress *in vitro*. Previous studies at the CBTBR showed that *M. smegmatis* log phase cultures treated with 2.5 mM H\(_2\)O\(_2\) resulted in a 4 log reduction in survival after 4 hours (Moolla, *et al.*, 2014; Hassim, *et al.*, 2015). Since H\(_2\)O\(_2\) is light sensitive and is unstable, variable results are obtained from one experiment to another hence, the optimal H\(_2\)O\(_2\) concentration to test the double DNA glycosylase deletion mutants generated in this study together with the previously generated single deletion mutants was optimized. *M. smegmatis* mc\(^2\)155 was grown to an OD\(_{600\text{nm}}\) of 0.35 and treated with 1.0, 2.0 and 2.5 mM of H\(_2\)O\(_2\). The data in Figure 23 indicate that 1.0 mM of H\(_2\)O\(_2\) was insufficient as
it did not display a killing effect after 6 hours of treatment. In contrast, at 2.5 mM all the cells
died after two hours of exposure to H₂O₂. Treatment of the bacteria with 2.0 Mm H₂O₂,
resulted in approximately 4-5 log killing after 4 hours of exposure to H₂O₂ hence, in this
study all further testing was done at 2.0 mM of H₂O₂.

**Figure 23.** Assessment of mc²155 with three different concentrations of H₂O₂. Early log phase early
log phase cultures were treated with 1.0 2.0 and 2.5 mM H₂O₂ and growth was monitored as CFUs/ml
over a 6 h period. The error bars represent the standard error between 3 biological replicates.

*M. smegmatis* mc²155, the double mutants and their complemented strains were grown to an
OD₆₀₀nm of 0.35 followed by treatment with 2.0 mM H₂O₂ and monitored for a period of 6
hours to determine whether the deletion of nth and mutY in each genetic background would
make the strains more prone to oxidative damage. The results from 5 independent
experiments showed some variation in the response of the wild type, double knockout
mutants and respective complemented strains under oxidative stress conditions (Figure 24
and Appendix 5.4, Figures 35 A-D). The variation amongst these experiments is likely linked
to the instability of H₂O₂. However, in three out of the five experiments (Figure 24 and
Appendix 5.4, Figures 35A and B) both the double deletion mutants showed a 2-3 log kill
after 4 hours of exposure to H₂O₂ which was exaggerated after 6 hours compared to the
parental strain. Previously, it was shown that the $\Delta\text{mutY}$ mutant was sensitive to oxidative stress (Hassim et al., 2015) and therefore, the observed phenotype of $\Delta\text{mutY}\Delta\text{nth::nth}$ complemented strain was as expected which was consistent with the previous observation by Hassim et al (Hassim et al., 2015). However, the $\Delta\text{mutY}\Delta\text{nth::nth}$ complemented strain displayed an increased sensitivity to oxidative stress after 4 hours (Figure 24 and Appendix 5.4 Figures 35A and B) which was surprisingly more than that observed for the double deletion mutants. The reason(s) for this observation is currently unclear. Previously, Moolla et al (Moolla et al., 2014) showed that loss of the nth gene does not affect survival under oxidative stress when compared to mc²155 and therefore, the $\Delta\text{nth}\Delta\text{mutY::mutY}$ complemented strain was expected to display $\text{H}_2\text{O}_2$ sensitivity at comparable to wild type levels. The reasons for the differences in these observations are currently unknown but the data clearly indicate that combinatorial loss of Nth together with MutY leads to increased mutagenesis and these two DNA glycosylases possibly co-ordinate their function to control mutagenesis in mycobacteria under oxidative stress conditions. In the other two experiments (Appendix 5, Section 5.4, Figures 35C and D) the double deletion mutant, $\Delta\text{mutY}\Delta\text{nth}$ and both the complemented strains showed the same kill kinetics as the parental strain indicating once again that some hierarchy exists between these DNA glycosylases as deletion of nth after the removal of mutY does not exaggerate mutagenesis. In both these experiments, the $\Delta\text{nth}\Delta\text{mutY}$ double deletion mutant surprisingly survived the oxidative assault which can be attributed to possible technical experimental error.
Figure 24. Survival of mc2155, ΔmutYΔnth, ΔnthΔmutY and complemented strains under oxidative stress condition. Early log phase early cultures were treated with 2.0 mM H\textsubscript{2}O\textsubscript{2} and growth was monitored as CFUs/ml over a 6 h period. The graph is a representative of one of five independent experiments. The other four representative graphs are shown in Appendix 5, Section 5.4.

3.5.4 H\textsubscript{2}O\textsubscript{2} diffusion susceptibility test

Since the H\textsubscript{2}O\textsubscript{2} liquid assay showed variability between experiments, the susceptibility test was performed on solid agar plates. The various mutants and respective parental strains were grown overnight to an OD\textsubscript{600nm} of 0.35 in 7H9 at 37 ºC. One Hundred microlitres of each culture was plated onto 7H10 plates and left at room temperature to allow the bacteria to be absorbed into the plate. Different concentrations of H\textsubscript{2}O\textsubscript{2}, 0.18, 0.35 and 0.44 mM (0.5, 1.0 and 1.25 µl) were spotted onto the centre of the 7H10 agar plates that had been inoculated with the test microorganism. Lower concentrations of H\textsubscript{2}O\textsubscript{2} were used in this assay as with 2.5 mM of H\textsubscript{2}O\textsubscript{2} there was complete killing (data not shown). Diffusion of H\textsubscript{2}O\textsubscript{2} in an outward manner from the spot creates a concentration gradient and after three days of incubation, the diameter of the zone of inhibition was measured. The control was 7H10 plates with the test organism without H\textsubscript{2}O\textsubscript{2} and the total numbers of cells (colony forming units CFU/ml) were
also calculated by spreading dilutions from the cultures onto 7H10. In this assay the loss of \( nth \) and \( mutY \) either individually or in combination did not significantly increase susceptibility to \( H_2O_2 \), compared to the wildtype strains (Figure 25). These data are in contrast to the liquid assay wherein the both the double mutants strains showed susceptibility to \( H_2O_2 \) compared to the mc\(^2\)155. It is possible that \( H_2O_2 \) in solid media is unable to generate sufficient oxidative stress to cause DNA damage resulting in variable kill kinetics in the two different assay systems.

![Graph of bacterial zones of inhibition](image)

**Figure 25.** \( H_2O_2 \) diffusion susceptibility assay of *M. smegmatis* and its derivative mutant and complemented strains. The graph represents the mean diameters of the bacterial zones of inhibition from three replicate experiments.

### 3.5.5 UV induced mutagenesis

Moolla *et al* (Moolla *et al.*, 2014) demonstrated that deletion of \( nth \) resulted in increased UV induced mutagenesis to rifampicin. Similarly, the *M. smegmatis* wildtype, single mutants, double mutants and the complemented strains generated in this study were tested for
enhanced UV induced mutagenesis. To determine the optimal UV energy that will result in mutagenesis, experiments were carried out using the wild type strain grown to an OD$_{600nm}$ of 0.6 and were treated with different UV fluences, 25, 50 and 70 mJ/cm$^2$ as described by (Boshoff et al., 2003) (Figures 26A and 26B). At all three wavelengths the cells displayed similar survival with about $1 \times 10^7$ cells recovered post UV exposure. There was a corresponding increase in rif$^R$ mutants at all three wavelengths as well and for all downstream experiments a UV fluence, of 50 mJ/cm$^2$ was used to induce mutagenesis.

**Figure 26.** Effect of different UV energies (25, 50, and 70 mJ/cm$^2$). (A) UV induced mutation frequency of *M. smegmatis* mc$^2$ 155 to rifampicin. (B) Survival of *M. smegmatis* mc$^2$155 after UV treatment.
All cultures were therefore, treated with 50 mJ/cm² and after treatment, cultures were inoculated into 50 ml 7H9 and allowed to recover for 3 and 6 hours before plating on 7H10. The single nth deletion mutant showed an increased mutation frequency to rifampicin resistance after 3 hours of recovery from UV in 7H9 compared to mc²155 (Figure 27) which is consistent with the observation of Moolla et al (Moolla et al., 2014), who demonstrated an increase in mutation frequency to rifampicin after 3 hours and marginal increase after 6 hours of recovery. In Figure 27, the ΔmutY deletion show a lower induced frequency of rifR compared to mc²155 and Δnth. It is possible that the intact nth gene can repair the damage caused by UV in this mutant strain or uses a different repair mechanism.

Figure 27. UV induced mutation frequency to rifampicin of mc² 155, Δnth, ΔmutY, Δnth::nth and ΔmutY::mutY. Results represent an average of three independent experiments.

Subsequently the mutation frequency to rifampicin resistance after UV induced damage was assessed for the two double deletion mutants, ΔmutYΔnth and ΔnthΔmutY to determine
whether combinatorial loss of two DNA glycosylases will enhance mutagenesis. The rif$^R$
 frequencies were not elevated when the double mutants were exposed to UV when compared
to mc$^2$155. However, in this instance the complemented ΔnthΔmutY::ΔmutY strain restored
phenotype of Δnth at 3 hours of recovery from UV (Figure 28). The ΔmutYΔnth::Δnth
complemented strain (Figure 28) did not restore the phenotype and the rif$^R$ mutation
frequency remained at the same level as the mutant strain. The elevated UV induced rif$^R$
frequency observed for the Δnth single deletion mutant and restoration of the phenotype in
the complemented ΔnthΔmutY::mutY strain at 3 hours of UV recovery suggests that nth is
clearly involved in the repair of UV induced DNA damage. These data are consistent with the
observation of Moolla et al (Moolla et al., 2014). Deletion of mutY did not show an
equivalent amount of mutagenesis as the nth deletion mutant (Figure 27) suggesting that
mutY is not involved in the repair of DNA damaged by UV. As the double mutants do not
show elevated UV induced mutagenesis, it is possible that other repair systems maybe
involved in maintaining mycobacterial genome integrity.
Figure 28. UV damaged induced mutation frequency of mc^2155, ΔmutYΔnth, ΔnthΔmutY, ΔmutYΔnth::nth and ΔnthΔmutY::mutY to rifampicin. Results represent an average of three independent experiments.

3.5.6 Mutation rates of Nth and MutY DNA glycosylases

Rates of mutation (μ) to rifampicin resistance were estimated by the fluctuation assay based on the use of the Luria and Delbruck analysis (Rosche and Foster, 2000). Mutation rate determinations are more accurate and reproducible compared to mutation frequency since they estimate the rate at which an organism will sustain a resistance confirming mutation to an antibiotic such as rifampicin during its lifetime. To investigate the combined role of Nth and MutY glycosylases on spontaneous mutation rates to rifampicin, the parental, Δnth, ΔmutY, ΔmutYΔnth, ΔnthΔmutY and the respective complemented strains were tested in triplicate in the fluctuation assay as described in the materials and methods (Section 2.6.6). Mutation rate calculations demonstrated (Table 3.1) that the individual loss of Nth or MutY did not result in an increased spontaneous mutation rate to rifampicin, which was in
concordance with two previous assessments of these mutant strains under these conditions (Moola et al. 2014; Hassim et al. 2015). However, in experiment 1 (Table 3.1; highlighted in blue) the nth deficient mutant strain displayed a 3.4 fold increase in mutation rate to rifampicin which could be attributed to technical differences and/or errors during the experimental procedure. Interestingly, the combined loss of Nth and MutY, regardless of the genetic order in which these respective genes were deleted, resulted in an overall increased mutation rate compared to the parental strain as well as the individual single deletion mutants ∆nth and ∆mutY, suggesting that these glycosylases play an important role in maintaining genome integrity in mycobacteria. However, the deletion of mutY in an nth deficient background, ∆nth∆mutY (highlighted in red) showed an exacerbated mutation rate compared to the ∆mutY∆nth (highlighted in orange) mutant wherein the nth was deleted in the ∆mutY mutant background. The ∆nth∆mutY mutant displayed between 2.2-2.5 fold differences in mutation rates compared to the ∆mutY∆nth mutant (Table 3.1). From these data it is clear that the two double mutants have different phenotypic characteristics that may be associated with genotypic changes in their respective genomes which may require whole genome sequencing to elucidate further. The complemented strains for the double deletion mutants showed variable levels of complementation between the three experiments, but in all cases, a reduction in mutation rate was observed. The lack of full complementation may be due to the fact that functional gene is not integrated in the exact genomic context (Table 3.1). From comparisons of the two double deletion mutants, based on the phenotypic analysis carried out in this study, it seems that the ∆nth∆mutY mutant has a less stable genome when compared to the ∆mutY∆nth mutant, as it consistently displayed a more exaggerated phenotype in response to oxidative stress and had a greater propensity to accumulate rifR mutations. This suggests that loss of mutY subsequent to nth seems to result in catastrophic lesions in M. smegmatis, which cannot be repaired by other DNA glycosylases of the base excision repair
pathway. The Nth glycosylase is also likely the more dominant of the two DNA glycosylases in controlling mutations under oxidative stress conditions.

Table 3.1: Spontaneous mutation rates to rifampicin of the single and double deletion mutants in the \(nth\) and \(mutY\) DNA glycosylases and their respective complemented strains.

<table>
<thead>
<tr>
<th>Stains</th>
<th>Experiment 1</th>
<th>Experiment 2</th>
<th>Experiment 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mutation rate</td>
<td>Fold increase</td>
<td>Mutation rate</td>
</tr>
<tr>
<td>mc(^2)155</td>
<td>4.21E-09</td>
<td>1</td>
<td>5.77E-09</td>
</tr>
<tr>
<td>(\Delta nth)</td>
<td>1.43E-08</td>
<td>3.40</td>
<td>5.40E-09</td>
</tr>
<tr>
<td>(\Delta mutY)</td>
<td>6.57E-09</td>
<td>1.56</td>
<td>2.03E-09</td>
</tr>
<tr>
<td>(\Delta mutY\Delta nth)</td>
<td>1.20E-08</td>
<td>2.9</td>
<td>1.03E-08</td>
</tr>
<tr>
<td>(\Delta nth\Delta mutY)</td>
<td>1.92E-08</td>
<td>4.6</td>
<td>4.37E-08</td>
</tr>
<tr>
<td>(\Delta nth::nth)</td>
<td>1.12E-08</td>
<td>2.66</td>
<td>4.21E-09</td>
</tr>
<tr>
<td>(\Delta mutY::mutY)</td>
<td>7.25E-09</td>
<td>1.72</td>
<td>2.94E-09</td>
</tr>
<tr>
<td>(\Delta mutY\Delta nth::nth)</td>
<td>1.10E-08</td>
<td>2.60</td>
<td>6.12E-09</td>
</tr>
<tr>
<td>(\Delta nth\Delta mutY::mutY)</td>
<td>1.19E-08</td>
<td>2.8</td>
<td>2.61E-08</td>
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</tbody>
</table>

3.5.7 Mutation spectrum

Rifampicin resistance arises mainly through the acquisition of mutations in the Rifampicin Resistance Determining Region (RRDR) within the 81-bp region of the \(rpoB\) gene (Figure 29) and the most common mutations occur at positions, 513, 522, 526, and 531 (Figure 29; highlighted in orange boxes). The RRDR region from genomic DNA isolated from 8-10 randomly selected rifampicin-resistant isolates from the fluctuation assay for mc\(^2\)155, \(\Delta nth\), \(\Delta mutY\), \(\Delta nth\Delta mutY\), \(\Delta mutY\Delta nth\) and their respective complemented strains was sequenced using the primer MsmrpoBF1 (Appendix 5.2, Table5.1). In the wildtype strain from 10
isolates 7 different mutations were revealed within the 81-bp region of \textit{rpoB} gene, with the highest percentage of mutations observed in \textit{His526 →Arg} (CAC → CGC) (30%), followed by \textit{Ser531 →Leu} (TCG → TGG) (Table 3.2). No mutations were found within the 81-bp \textit{rpoB} segment for two rif\textsuperscript{R} isolates, presumably the mutations occurred outside the sequenced region. In the \textit{Δnth} mutant there was an increase in C→T mutations at \textit{Ser531 → Leu} (TCG → TTG) from 20\% in mc\textsuperscript{2}155 to 30\%. Also, there was an increase in mutation at \textit{Ser531 → Trp} (TCG → TGG) from 10\% in the mc\textsuperscript{2}155 wildtype to 20\% in the mutant (Table 3.2). In a previous study within the laboratory, the \textit{ΔmutY} mutant spectrum displayed increased C→A transversions at Gln513 → Lys (CAG → AGG) (Hassim \textit{et al.}, 2015). Similarly in this study \textit{ΔmutY} mutant yielded in increased C→A transversions (44.4\%) at the same position. This phenotype was reversed in the complemented strain (Table 3.2).

Interestingly, the deletion of \textit{nth} in \textit{ΔmutY} (\textit{ΔmutYΔnth}) yielded in the highest mutation frequency (33\%) at the Asp516→Tyr (GAC → TAC). This point mutation was unique to this mutant and was absent in the other complemented strains (Table 3.2). The prevalence of the Asp516 substitution in \textit{ΔmutYΔnth} was unexpected because the mutation occurred in the amino acid which is not common within the RRDR region. The deletion of \textit{mutY} in \textit{Δnth} (\textit{ΔnthΔmutY}) presented an increase in \textit{His526 → Tyr} mutations and this was in concordance with the increased C→T transitions (Ser531→Leu), which also formed the major mutations in the \textit{nth} single deletion mutant (Table 3.2). There was also an increase of 20\% in C → A mutations at Gln513 → Lys (CAG→AGG) when compared to mc\textsuperscript{2}155. The increase in C→T and C→A transitions in the \textit{ΔnthΔmutY} was expected as the Nth and MutY DNA glycosylases are responsible for repairing these mutational changes.
**Figure 29.** The 81 bp RRDR region of the *rpoB* gene. The common areas where mutations occur are highlighted in orange boxes.
Table: 3.2. Mutation patterns and percentage distribution of the rifampicin resistant mutations found in the RRDR region. The total number of sequenced rifR mutants for each strain is shown in brackets. The common amino acids where mutations occur are in blue and the least common are in black. Values are given as a percentage.

<table>
<thead>
<tr>
<th>Mutation</th>
<th>Amino acid change</th>
<th>mc² 155(10)</th>
<th>∆nth (10)</th>
<th>∆mutY (9)</th>
<th>∆mutY ∆nth (9)</th>
<th>∆nth ∆mutY (10)</th>
<th>∆nth ::nth (8)</th>
<th>∆mutY ::mutY (9)</th>
<th>∆mutY ∆nth ::nth (10)</th>
<th>∆nth ∆mutY ::mutY (10)</th>
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</thead>
<tbody>
<tr>
<td>CAC → GAC</td>
<td>His526 → Asp</td>
<td>10</td>
<td>-</td>
<td>-</td>
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<td>-</td>
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<td>-</td>
<td>-</td>
</tr>
<tr>
<td>TCG → TGG</td>
<td>Ser531 → Trp</td>
<td>10</td>
<td>20</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CAC → TAC</td>
<td>His526 → Ser</td>
<td>10</td>
<td>-</td>
<td>-</td>
<td>10</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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</tr>
<tr>
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<td>His526 → Arg</td>
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<td>10</td>
<td>11.1</td>
<td>22.2</td>
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<tr>
<td>CGT → TGT</td>
<td>Leu 521 → Cys</td>
<td>-</td>
<td>-</td>
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<td>-</td>
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<td>His526 → Tyr</td>
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<td>Arg528 → His</td>
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<td>-</td>
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<td>62.5</td>
<td>11.1</td>
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<td>CTG → GTG</td>
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<td>CAC → ACC</td>
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<td>CAG → CGG</td>
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<tr>
<td>GGT → GAT</td>
<td>Gln513 → Asp</td>
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<td>10</td>
</tr>
<tr>
<td>GAC → TAC</td>
<td>Asp516 → Tyr</td>
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<td>-</td>
<td>-</td>
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<td>-</td>
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<td>CGT → TGT</td>
<td>Arg529 → Cys</td>
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<td>-</td>
<td>-</td>
<td>25</td>
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4.0 Discussion

The ability to rapidly repair DNA damage caused by reactive oxygen and nitrogen intermediates in the host environment using robust DNA repair systems is essential for \textit{M. tuberculosis} to survive in hostile conditions within phagocytic cells. Consequently, to counter these threats \textit{M. tuberculosis} is armed with efficient repair systems such as base excision repair (BER) and the GO repair system. The key enzymes within the BER pathway are the DNA glycosylases. Our interest in these repair pathways stemmed from a need to understand the importance and functioning of the DNA glycosylase enzymes in mycobacteria, more specifically, the role of Nth, Fpg, Nei and MutY in controlling mutations within this species. A key question which remained unanswered is whether genes involved in BER work independently or interdependently. According to previous findings in our laboratory, there is strong evidence of overlapping functions between these DNA glycosylase genes as well as a backup function in the event of extensive DNA damage. The study by Moolla \textit{et al} demonstrated that the Nth and Nei DNA glycosylases functioned synergistically and probably compensate for each other under oxidative stress conditions in order to maintain genomic integrity (Moolla \textit{et al}., 2014). Similarly, the loss of MutY together with the Fpg DNA glycosylases displayed an exaggerated reduction in survival under oxidative stress conditions and an increased spontaneous mutation rate to rifampicin, suggesting a co-operative interaction between these two proteins for genome maintenance (Hassim \textit{et al}., 2015). These findings highlighted that Nth and MutY have an antimutator role and the Fpg/Nei family of DNA glycosylases play more of a backup role during DNA repair under oxidative stress conditions. Hence, in this study, we attempted to understand if there was functional association or overlap between the Nth and MutY DNA glycosylases in the BER pathway. To our knowledge the combined loss of these DNA glycosylases for the maintenance of genome
integrity under stressful conditions has not been previously investigated in *E.coli* or in any mycobacterial species.

The Nth and MutY glycosylases are members of a sequence related family involved in the BER pathway with MutY forming part of the GO system. Panels of single DNA glycosylase gene knockout mutants were available from previous studies (Moolla *et al*., 2014, Hassim *et al*., 2015) hence, these were used to generate the two double mutants that lacked both Nth and MutY in different genetic lineages. The *mutY* gene was deleted in a mutant strain that had the *nth* gene deletion (Moolla *et al*., 2014) and in the second double mutant *nth* was deleted in a *mutY* deletion mutant that was previously generated (Hassim *et al*., 2015). In this study, the individual and combined roles of the *M. smegmatis* Nth and MutY DNA glycosylases was assessed under normal growth culture conditions and under different stress conditions to determine whether the loss of these enzymes had an impact on survival of the mutant strains. Loss of key DNA repair enzymes is assumed to increase mutations hence, the changes in spontaneous mutation rates to rifampicin was measured for each of the combinatorial deletion mutants. Collectively, the data presented here have afforded greater insight into the functioning of the BER pathway in mycobacteria and clearly show that both Nth and MutY are required to prevent mutations in the mycobacterial genome.

As previously shown (Moolla *et al*., 2014, Hassim *et al*., 2015), the single *nth* or *mutY* deletion mutants which were used as control strains in this study showed the same *in vitro* growth kinetics as the parental strain under normal culture conditions. The combined loss of these DNA glycosylases did not affect growth as both the double mutants displayed the same doubling times as the single deletion mutants and the parental wild type strain, mc²155. These data suggest that the two glycosylases are not essential for growth under *in vitro* culture
conditions or that there is not enough spontaneous DNA damage generated under these conditions to cause growth defects.

In previous studies, 2.5 mM H$_2$O$_2$ was used to generate *in vitro* oxidative stress conditions however, in this study under these conditions this led to complete killing of the parental strain after 2 hours of exposure. Hence, the H$_2$O$_2$ concentration was optimized, and with 2.0 mM H$_2$O$_2$ a 5 log reduction (10$^3$ viable cells) was observed for the wildtype strain after 4 hours of exposure. Subsequently, assessment of the parental mc$^2$155, the double mutant strains and their respective complemented strains were tested with 2.0 mM H$_2$O$_2$ and the data revealed variable killing rates between biological repeat experiments. This discrepancy between experiments could be attributed to the instability of H$_2$O$_2$, which is affected by a range of external sources, one of which is light. Interestingly, based on the 5 biological repeats there was a consistent trend of almost equivalent decreased survival for both the double deletion mutants after 6 hours of exposure to H$_2$O$_2$ compared to mc$^2$155 in three of the experiments. However, the complemented strains were unable to restore the defect and in fact showed a greater reduction in survival compared to the mutant strains. In two of the experiments the ΔmutYΔnth mutant showed enhanced survival under oxidative stress suggesting that there may be other protective mechanisms that may have been triggered in this mutant to allow it to combat the effects of oxidative stress. Due of the variation in biological replicates in the liquid assay, we employed a more stable H$_2$O$_2$ spotting assay to test for sensitivity of the strains. In contradiction to the trend we saw in the liquid assay of H$_2$O$_2$, the ΔmutYΔnth mutant showed no difference in survival when compared to the single mutants or mc$^2$155 wildtype. The differences in the results might be due to H$_2$O$_2$ being more effective in the liquid assay in comparison to the spotting assay since the strains take approximately 3-5 days
to grow on the plate and by this time the H₂O₂ could lose its effectiveness as it is light sensitive.

The phenotypic differences between the two double mutants (AmutYΔnth and ΔnthAmutY) could be as a result of a compensatory mutation that may have been introduced during the generation of the single mutY mutant. In order to understand the phenotypic differences manifested between these two double mutant strains from different genetic lineages, whole genome sequencing will be required, however that is beyond the scope of this project.

In E.coli, the involvement of Nth in the repair of DNA damage caused by UV has long been suspected since it was initially discovered using UV-irradiated DNA substrates (Radman, 1976). The Nth protein in E.coli is a key enzyme responsible for the removal of oxidized pyrimidines (Bauch and Laval, 1999, Zharkov and Grollman, 2002). To determine the exact role of Nth in the repair of UV induced lesions in mycobacteria, Moolla et al. (Moolla et al., 2014) assessed the Δnth single mutant for UV (250 mJ/cm²) induced mutagenesis. The single Δnth mutant, 3 hours post UV treatment, demonstrated increased mutation frequencies compared to the parental mc²155 strain, suggesting its role in controlling UV induced mutagenesis. Similarly, in this study the single deletion mutants of nth together with the two combinatorial double deletion mutants, were tested for increased mutagenesis under UV (50 mJ/cm²) induced conditions. The data for the single nth deletion mutant showed an almost 10 fold increase in mutagenesis compared to the parental strain after 3 hours of recovery post UV which was in accordance to previously observed data (Moolla et al., 2014). However, there was only a marginal change in mutation frequency for the nth deletion mutant after 6 hours, which was not significantly different to the parental strain. Moreover, the complemented strain ΔnthΔmutY::mutY when exposed to UV showed the same mutation.
frequencies as the single \textit{nth} mutant confirming that Nth plays an important role in repairing UV induced DNA damage. In contrast the \textit{mutY} single deletion mutant or either of the two double deletion mutants showed no increased mutagenesis after 3 and 6 hours post UV treatment compared to the parental strain. A likely explanation for the reduced induced frequencies could be the suppression of UV induced mutations by \textit{mutY} or its non-involvement in the repair of damage caused by UV. Consistent with this hypothesis is the observation of no change in UV induced mutagenesis in a \textit{Helicobacter pylori} \textit{mutY} deficient mutant compared to the parental strain, suggesting that MutY may not play a role in UV induced mutagenesis (Huang \textit{et al.}, 2006).

A compromised DNA repair system impacts on the integrity of the genome resulting in increased mutations. The various single and double mutants (mc\textsuperscript{2}155, \Delta\textit{nth}, \Delta\textit{mutY} and \Delta\textit{mutY}\Delta\textit{nth}, \Delta\textit{nth}\Delta\textit{mutY}) were assessed for changes in spontaneous mutation frequency and rates to rifampicin. The single \textit{nth} or \textit{mutY} deletion mutants and the \Delta\textit{mutY}\Delta\textit{nth} double mutants yielded a two fold increase in spontaneous mutation frequency when compared to mc\textsuperscript{2}155 wildtype. In contrast, the double mutant strain (\Delta\textit{nth}\Delta\textit{mutY}), that showed an enhanced survival trend under oxidative stress conditions also showed a significant elevation in (3 fold higher) in spontaneous rifampicin mutagenesis compared to the single deletion mutants and the double mutant \Delta\textit{mutY}\Delta\textit{nth}. The striking difference in mutation frequency between the two double mutants is somewhat surprising given the fact that both these strains are devoid of the same DNA glycosylases and therefore, should be genetically identical. The only difference is the order in which they were inactivated and thus it is highly likely that during the generation of the \Delta\textit{mutY}\Delta\textit{nth} double mutant, compensatory mutation(s) may have been introduced within the genome, which may have imparted a fitness advantage to this mutant. In previous studies, the loss of \textit{mutY} did not show an increase in mutation rates.
(Kurthkoti et al., 2010, Hassim et al., 2015) however, reassessment of the mutY deletion mutant in this study, revealed a two-fold higher mutation frequency when compared to the wildtype. The variation between these experiments might be due to mutations occurring early during culture growth, leading to the propagation of the same mutation during the life-cycle, a phenomenon known as a “jackpot phenotype”(Luria and Delbrück, 1943). The mutation frequency data for the various mutants and parental strain was further confirmed with mutation rate measurements using the Luria-Delbrück fluctuation assay. Mutation rate determinations are more accurate and reproducible compared to mutation frequency as the estimate the rate at which an organism will sustain a resistance mutation to an antibiotic during its lifetime. In contrast to the spontaneous mutation frequency, assessment of spontaneous mutation rates to rifampicin showed no increase in mutation rate for the single Δnth and ΔmutY mutants compared to mc²155 which is consistent with previous findings (Moolla et al., 2014, Hassim et al., 2015). Once again the ΔnthΔmutY double mutant strain showed a dramatic increase (4.6 to 7.6 fold) in mutation rate but a comparable significant increase was not observed for the ΔmutYΔnth mutant (1.8 to 3.4 fold). The collective data obtained in this study clearly shows that Nth could be classified to have a more important an antimutator role for genome stability in mycobacteria compared to MutY.

More than 90% of rifampicin resistance in M. tuberculosis strains is due to mutations acquired within an 81 bp region within the rpoB gene (Ramaswamy and Musser, 1998). The mutations consist of nucleotide changes, deletions and insertions (Yue et al., 2003) In mycobacteria the most common codons which sustain mutations are at position 512,522, 526 and 531 (Höfling et al., 2005, Hillemann et al., 2005). In order to identify the mutational changes responsible for rifampicin resistance the 81 bp RRDR region was sequenced for several rifampicin resistant colonies obtained for mc²155 wildtype, the single mutants, double
mutants and their respective complemented strains from the fluctuation assay. Previous studies showed that deletion of mutY leads to increased C → A mutations in M. smegmatis (Kurthkoti et al., 2010, Hassim et al., 2015). Similarly, in this study the mutY deletion mutant showed a 34% increase in C → A mutations compared to the mc2155 wildtype strain. Since damaged pyrimidine bases and oxidised cytosines are generally recognized by the Nth glycosylase, in the absence of this enzyme it is expected that there will be an increase in the C→T or G→A mutations (Guo et al., 2010). Consistent with this, in the single nth deletion mutant the majority of mutations were C→T changes, with a combined total of 40%. In the double mutants, which are devoid of the nth and mutY DNA glycosylases, an increase in C → A and A → C mutations attributed to the deletion of mutY, and C→T or G→A mutations due to the absence of nth was as expected. Interestingly, in the ΔmutYΔnth mutant there was a unique mutation at codon 516 (G → T), which was not present in any of the strains analysed in this study. The ΔnthΔmutY strain showed an increased frequency of C→T transitions compared to the wildtype but this was comparable to the single nth mutant. There was also a marginal increase in C → A transversions which could be attributed to the loss of mutY. The two double mutants (ΔmutYΔnth and ΔnthΔmutY) also displayed a diverse spectrum of rifR mutations once, again suggesting that the two mutants generated from different lineages have genetic variances.

In conclusion, this study further highlights key insight regarding the importance of Nth and MutY DNA glycosylases in the BER pathway in maintaining genome integrity. Based on the data obtained, it is clear that the combinatorial loss of Nth and MutY does have an additive impact on the survival of M. smegmatis under oxidative stress conditions, compared to the individual enzymes respectively. Based on the phenotypic differences observed for the two double mutants, it is possible that initial loss of nth which has been shown to have
antimutator properties (Moolla et al., 2014) may be predisposing the mycobacteria to catastrophic levels of DNA damage hence, subsequent deletion of *mutY* exaggerates genomic instability. As *mutY* does not have scission activity removal, of MutY first does not have a severe genotoxic impact on the genomic DNA as the remaining Fpg/Nei and Nth DNA glycosylases are able to act on any mutations arising under oxidative stress. Moreover, these results may suggest a functional hierarchy between the various DNA glycosylases in the BER system with Nth predominating over MutY as an antimutator.

4.1 Future studies

As the oxidative stress inducing agent H$_2$O$_2$ used in this study had limitations, it is important that future studies focus on testing and optimizing more stable compound for generating oxidative stress. Testing the panel of deletion mutants under improved conditions may shed further insight on the association and interplay of these various DNA glycosylases during DNA repair. In the absence of Nth and MutY it would also be interesting to measure the expression levels of Fpg and Nei under normal culture conditions as well as under oxidative stress conditions to assess whether these DNA glycosylases display a backup function. Whole genome sequencing of the two double mutants will identify genomic differences in terms of compensatory mutations that may be contributing to the phenotypic difference observed for the two strains. In addition, perhaps other DNA repair genes in the BER pathway may be identified to play a role in genome maintenance.
5. Appendix

5.1 Media preparation and solutions

5.1.1 *Mycobacterium smegmatis*

**Middlebrook 7H9**

4.9 g powder, 0.2% glycerol, 0.05% Tween 80 (filter sterilized) and 0.2% glucose

**Middlebrook 7H10**

19 g powder, 0.085 % NaCl, 0.2% glucose, 0.2% glycerol

5.1.2 *Escherichia coli*

**Luria-Bertani Broth (LB)**

5g yeast extract, 10g tryptone, 10g NaCl.

All media were made up to 1000 ml and autoclaved at 121°C for 10 min.
## 5.2 PCR and sequencing primers

**Table 5.1.** Primers used in this study, with description and expected sizes.

<table>
<thead>
<tr>
<th>Oligonucleotide name</th>
<th>Sequence 5’ – 3’</th>
<th>Description</th>
<th>Expected (bp)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>NthF1, NthR1</td>
<td>GTCAGCCGTGGTCAGT ACC\nCGAGACGCTCATCGTG TG</td>
<td>Confirmation of wildtype and DCO</td>
<td>Wildtype (833)\nDCO (270)</td>
<td>(Moolla et al., 2014)</td>
</tr>
<tr>
<td>mutYFwd, mutYRev</td>
<td>GCCCTGAATTCTGCTGA CAC\nCGTGACACCCTCGGAC TAC</td>
<td>Confirmation of wildtype and DCO</td>
<td>Wildtype (1481)\nDCO (780)</td>
<td>(Hassim et al., 2015)</td>
</tr>
<tr>
<td>NthUsF1, NthUsR1</td>
<td>AAGCTTGTGTGCGTGCT GGCCAAG\nAGATCTCACACCGGTG ATCGTCTT</td>
<td>Nth upstream deletion region</td>
<td>1185</td>
<td>(Moolla et al., 2014)</td>
</tr>
<tr>
<td>NthDsF2, NthDsR2</td>
<td>AGATCTACGACGCAGCAG\nGGTACCAGCTGGGGCA TCTCTTCC</td>
<td>Nth Downstream deletion region</td>
<td>1170</td>
<td>(Moolla et al., 2014)</td>
</tr>
<tr>
<td>MutYUSF, MutYUSR</td>
<td>GGCCAAGCTTAGCTGA TCGTTCGCGACTAC\nGCCAGATCTCGCATGG CTGCCAAGAGGTTA</td>
<td>mutY Downstream deletion region</td>
<td>1932</td>
<td>(Hassim et al., 2015)</td>
</tr>
<tr>
<td>MutYDSF, MutYDSR</td>
<td>GGCCAAGCTCGACTCG CTGCTGGTGAGC\nGCCGCCCGCCGCGCTT CTGCCGACAGAGGTTA</td>
<td>mutY Downstream deletion region</td>
<td>1775</td>
<td>(Hassim et al., 2015)</td>
</tr>
<tr>
<td>MsmrpoBF2, MsmrpoBR2</td>
<td>TGCGTGCCTGCACCGA GGTC\nATCTGTCCGGTGACCA CACCG</td>
<td>RRDR region</td>
<td>549</td>
<td>(Hassim et al., 2015)</td>
</tr>
<tr>
<td>MsmrpoBF1</td>
<td>GCAGACCTGATCAAC ATCC</td>
<td>RRDR sequencing primer</td>
<td>451</td>
<td>(Hassim et al., 2015)</td>
</tr>
</tbody>
</table>
5.3 Vectors maps used in this study and previous studies

Figure 30 GEM3Zf(+). Vector used for cloning upstream and downstream fragments to generate the \nth and \mutY deletion knockout constructs (Moolla \textit{et al.}, 2014 and Hassim \textit{et al.}, 2015). The plasmid was also used as a positive control for \textit{E.coli} transformations.
**Figure 31.** p2NIL suicide vector The suicide vector used in 3 way cloning, where the upstream and downstream fragments for the *nth* and *mutY* deletion constructs were simultaneously ligated into p2NIL (Parish and Stoker, 2000).
Figure 32. pGOAL17. The vector with the Pac cassette (sacB, lacZ,) which was inserted into p2NIL vector carrying the mutY deletion allele to generate p2NILΔmutY::pGOAL17 (Hassim et al., 2015).

Figure 33. pGOAL19. The vector with the Pac cassette (sacB, lacZ, and hyg) which was inserted into the p2NIL vector carrying the nth deletion allele to generate p2NILΔnth::pGOAL19 (Moolla et al., 2014).
**Figure 34.** pTWEETY. Vector used for constructing the all the complementation vectors.

### 5.4: Effect of hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}) on the survival of *M. Smegmatis*

![Graph showing survival of different strains of M. Smegmatis over exposure to H\textsubscript{2}O\textsubscript{2}](image)
**Figure 35.** Survival of the mc\(^2\)155, ΔmutYΔnth, ΔmutYΔnth and complemented strains under oxidative stress conditions. Early log phase cultures were treated with 2.0 mM H\(_2\)O\(_2\) and growth was monitored as CFUs/ml over a 6 h period.

5.5 DNA molecular weight markers

**Figure 36.** The DNA molecular weight markers. A. Marker λIV. B. Marker λVI used in this study (Roche Biochemicals)
6. References


CADET, J. & WAGNER, J. R. (2014) Oxidatively generated base damage to cellular DNA by hydroxyl radical and one-electron oxidants: Similarities and differences. *Archives of biochemistry and biophysics*, 557, 47-54.


