

Sample Honours Thesis Table of Contents

Please note: this is only an **example**. Each School has its own specifications, some of which are stricter than others. Furthermore, each thesis is different and will have different emphases on particular sections. CHECK with your supervisor for advice on length of sections and of the thesis as a whole.

Sample: *from the School of* BABS, UNSW

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(TAN, 2004,PP.II-IV)