BMJ Open

Examining the predictors of academic outcomes for indigenous Māori, Pacific and rural students admitted into medicine via two equity pathways: a retrospective observational study at the University of Auckland, Aotearoa New Zealand

Elana Curtis,1 Erena Wikaire,1 Yannan Jiang,2 Louise McMillan,2 Robert Loto,1 Phillippa Poole,3 Mark Barrow,4 Warwick Bagg,5 Papaarangi Reid1

ABSTRACT

Objective To determine associations between admission markers of socioeconomic status, transitioning, bridging programme attendance and prior academic preparation on academic outcomes for indigenous Māori, Pacific and rural students admitted into medicine under access pathways designed to widen participation. Findings were compared with students admitted via the general (usual) admission pathway.

Design Retrospective observational study using secondary data.

Setting 6-year medical programme (MBChB), University of Auckland, Aotearoa New Zealand. Students are selected and admitted into Year 2 following a first year (undergraduate) or prior degree (graduate).

Participants 1676 domestic students admitted into Year 2 between 2002 and 2012 via three pathways: GENERAL admission (1167), Māori and Pacific Admission Scheme—MAPAS (317) or Rural Origin Medical Preferential Entry—ROMPE (192). Of these, 1082 students completed the programme in the study period.

Main outcome measures Graduated from medical programme (yes/no), academic scores in Years 2–3 (Grade Point Average (GPA), scored 0–9).

Results 735/778 (95%) of GENERAL, 111/121 (92%) of ROMPE and 146/183 (80%) of MAPAS students graduated from intended programme. The graduation rate was significantly lower in the MAPAS students (p<0.0001). The average Year 2–3 GPA was 6.35 (SD 1.52) for GENERAL, which was higher than 5.82 (SD 1.65, p=0.0013) for ROMPE and 4.33 (SD 1.56, p<0.0001) for MAPAS. Multiple regression analyses identified three key predictors of better academic outcomes: bridging programme attendance, admission as an undergraduate and admission GPA/Grade Point Equivalent (GPE). Attending local urban schools and higher school deciles were also associated with a greater likelihood of graduation. All regression models have controlled for predefined baseline confounders (gender, age and year of admission).

Conclusions There were varied associations between admission variables and academic outcomes across the three admission pathways. Equity-targeted admission programmes inclusive of variations in academic threshold for entry may support a widening participation agenda, however, additional academic and pastoral supports are recommended.

INTRODUCTION

Widening participation in the medical profession remains a priority for many countries worldwide.1 2 Most medical schools acknowledge the need to embrace a widening participation agenda in order to contribute to the development of a health workforce that reflects a community’s ethnic, cultural,
geographic and socioeconomic diversity. Health workforce diversity is expected to reduce inequities in health outcomes through enhanced patient–provider interactions, increased provision of culturally competent care and better delivery to high-need, underserved population groups. In addition to workforce and healthcare delivery benefits, increasing diversity within medical school classes has been associated with positive effects on the medical school context itself including enhanced educational experiences for all students, positive student attitudes towards the value of diversity within medicine and the creation of learning contexts that challenge stereotypes and reduce implicit bias of medical students towards under-represented minorities. Widening participation interventions have been successful at increasing medical school diversity for under-represented ethnic minorities, women and rural students; however, disparities by socioeconomic status remain, as reported in the UK. Despite the strong rationale and increasing evidence of effectiveness, interventions to widen participation, such as medical school quotas, regularly come under attack and are criticised for lowering academic and quality standards. Comprehensive data analyses that measure outcome differences by admission pathways and attempt to examine the likely predictors for any observed differences are needed. This information is expected to better inform the widening participation debate and assist institutions to provide appropriate recruitment and tertiary support interventions for students admitted under equity-targeted admission pathways.

This study explores the predictors of both short-term and long-term academic outcomes for (1) indigenous Māori or Pacific students and (2) rural background students admitted into the medical programme (MBChB) under equity admission pathways, compared with general admission at the Faculty of Medical and Health Sciences (FMHS), University of Auckland (UoA) and Aotearoa New Zealand (NZ). This is one of two medical schools in NZ, based in a city of over 1.2 million, about one-third of the nation’s population. Entry into the MBChB at UoA may occur in two ways as: (1) an undergraduate within the first year of a health sciences or biomedical sciences degree at the UoA or (2) a graduate with a completed undergraduate or postgraduate qualification. Both pathways equate to ‘Year 1’ of the MBChB degree at the UoA. The Māori and Pacific Admission Scheme (MAPAS) commenced in 1972 in response to Māori and Pacific health workforce shortages, significant inequities in health outcomes and the indigenous rights of Māori within NZ. MAPAS involves comprehensive recruitment and retention interventions delivered within culturally appropriate contexts of support with approximately 240 MAPAS medical students enrolled in 2017 (approximately 20% of the total cohort). The Rural Origin Medical Preferential Entry (ROMPE) pathway began in 2004, in response to NZ government prioritisation of rural healthcare needs and evidence that students from rural backgrounds are more likely to return to practice in rural regions.

Over the first 20 years of MAPAS (ie, 1972–1992), there was a higher withdrawal rate for MAPAS medical students compared with other students admitted; however, the reasons for these findings are unclear and no associations between likely predictor variables and academic outcomes have been investigated to date. We hypothesise that markers of socioeconomic status, transition factors, bridging programme attendance (implemented specifically for Māori and Pacific students aspiring to enter medicine from 1999) and prior academic preparation, are likely to impact on both short-term, that is, Year 2–3 Grade Point Average (GPA) and long-term, that is, graduation outcomes. This study aimed to examine the association between admission variables and academic outcomes for students admitted into the medical programme under equity admission pathways in comparison to those students admitted under the general (usual) admission pathway.

METHODS

Study design

A retrospective observational study design was used to analyse data from all domestic students entering Year 2 MBChB at the UoA between 2002 and 2012 (with graduation data inclusive of academic outcomes from 2013). International students were excluded from analysis. Individual student demographic, admission and academic results data were sourced from Student Services Online, the UoA’s web-based centralised student data management system and the Medical Programme Directorate within the FMHS. The study period reflects the availability of electronic data from these sources and the time required for students to have graduated from a 6-year medical programme at the time this study commenced. A Kaupapa Māori Research (KMR) framework, supplemented by Pacific research methodology, was used throughout all aspects including study design, data collection, data analysis and research dissemination. This approach includes: a commitment to ensuring that the research outputs will have positive benefits for Māori and Pacific participants and communities; an explicit challenge to reject ‘victim blame’ and ‘cultural deficit’ analyses when interpreting data; and ensuring that any recommendations made from the research aim to facilitate participant academic success. This broad approach...
is expected to provide benefit for all study participants. The study was approved by the UoA Human Participants Ethics Committee (UAHPEC) (Reference 8110).

**Predictor variables**
Participants were identified by their admission category (MAPAS, ROMPE, GENERAL). The decile rating of secondary school attended was used as a proxy measure of socioeconomic status: low (1–3) (high deprivation), medium (4–7) and high (8–10) (low deprivation). High decile schools have a high proportion of students who reside in areas of low deprivation (high socioeconomic status). Attended school in Auckland (yes, no) and admitted into Year 1 as a school leaver (SL; yes, no) were used to measure transitioning effects, that is, impact of relocation to Auckland City (the largest city in NZ with a population of 1.4 million where the UoA medical programme is based) and impact of beginning tertiary study as a mature student or SL entrant. SL is defined as enrolment in bachelor level study in the year immediately following secondary school. Completion of a UoA bridging foundation programme (yes, no) that aims to bridge the ‘gaps’ between secondary and tertiary education contexts was recorded. The entry pathway into Year 2 MBChB was recorded as graduate or undergraduate. Academic preparation for medical entry was measured by the GPA or Grade Point Equivalent (GPE) at the time of admission for undergraduate and graduate applicants, respectively (0–9 representing Fail to A+ average grade).

**Outcome variables**
Two outcome variables were included in this study: graduated from MBChB (yes, no) and MBChB Year 2–3 GPA (0–9). Graduated from MBChB represents a long-term academic outcome and was only applied to those students who completed the MBChB programme by 2013, that is, students admitted between 2002 and 2009. The Year 2–3 GPA represents a short-term academic outcome associated with the 2 preclinical years of the MBChB programme. Data for this measure were available for a larger cohort of current and graduated students, that is, students admitted between 2002 and 2012. The Year 2–3GPA represents the average GPA achieved across Years 2 and 3 for students admitted between 2002 and 2011, and the GPA achieved across Year 2 only for students admitted in 2012.

**Analysis**
Statistical analysis was performed using SAS version 9.4. All statistical tests were two sided at a 5% significance level. A full statistical analysis plan was developed a priori that incorporated baseline confounders, key predictor and outcome variables of interest, based on concepts identified from relevant health workforce development and tertiary education literature as well as experience within the FMHS context as to the factors likely to impact on student success (figure 1). Multiple regression analyses with stepwise model selection were used to test the associations between predictor variables and academic outcomes for the total cohort (ie, MAPAS, ROMPE and GENERAL admission combined) and via entry admission subcohorts (ie, MAPAS and GENERAL). The results on ROMPE were not included due to small number of students in the study cohort.

The baseline model has controlled for predefined confounders including gender, age and year of admission into Year 2 MBChB (model 1) with the addition of predictor variables representing the sequential effect of socioeconomic status (model 2), transitioning (model 3), bridging programme (model 4) and academic preparation (model 5) on academic outcomes. Each model was initially run with all the prespecified predictors of interest, and those predictors that were significant at the 5% level were retained in the final model. This analysis was applied to all students admitted under MAPAS, ROMPE and GENERAL categories, with the outcome variables assessed at the time of data collection. For MBChB Year 2–3GPA, the mean difference was reported with
95% CI using the linear regression model. For graduation outcome (yes/no), the OR was reported with 95% CI using the logistic regression model. Similar regression analyses were conducted on the two largest subcohorts for MAPAS and GENERAL categories separately, in order to identify significant predictors of academic outcomes specific to that subcohort. Ethnicity was added to the baseline model in the subcohort analyses for MAPAS (Māori, Pacific) and GENERAL (Māori, Pacific, Asian, European/Pākehā, other/missing). Missing data were reported in the descriptive summary, but excluded in final regression analysis.

RESULTS
A total of 1676 students were included in the study, representing 1167 (70%) GENERAL, 317 (19%) MAPAS and 192 (11%) ROMPE admission categories. Cohort demographics are presented in table 1.

The MAPAS category differs in comparison to the GENERAL category by ethnicity (59% Māori, 41% Pacific, 0.3% Asian, 0.3% other, p<0.0001), school decile (42% high, 34% medium and 20% low, p<0.0001), having attended an Auckland school (60%, p<0.0001) and being admitted into medicine as an SL (50%, p<0.0001). The average admission GPA/GPE was approximately two points lower for MAPAS compared with GENERAL admission category students (6.22, SD 1.19, p<0.0001). The ROMPE category differs in comparison to the GENERAL category by mean age (21.5, SD 4.55, p<0.0001), gender (61.5% female, p<0.017), ethnicity (84% European/Pākehā, 11% Asian, 1% Māori, 1% Pacific, 2% other, p<0.0001), school decile (43% high, 43% medium, 7% low, p<0.0001), having attended an Auckland school

Table 1 Descriptive variables for GENERAL, MAPAS and ROMPE admission categories 2002–2009

| Descriptive summary variables | GENERAL (n=778) | MAPAS (n=183) | p-Value | ROMPE (n=121) | p-Value | Total (N=1082) | p-Value |
|-----------------------------|----------------|--------------|---------|---------------|---------|----------------|---------|
| Categorical variables | n | % | n | % | p-Value | n | % | p-Value | n | % |
| Female | 427 | 54.88 | 103 | 56.28 | 0.7319 | 76 | 62.81 | 0.1023 | 606 | 56.01 |
| Ethnicity | | | | | | | | | | <0.0001 |
| Māori | 20 | 2.57 | 102 | 55.74 | <0.0001 | 2 | 1.65 | <0.0001 | 124 | 11.46 |
| Pacific | 14 | 1.80 | 79 | 43.17 | <0.0001 | 2 | 1.65 | 95 | 8.78 |
| Asian | 339 | 43.57 | 1 | 0.55 | <0.0001 | 12 | 9.92 | 352 | 32.53 |
| Other | 44 | 5.66 | 1 | 0.55 | <0.0001 | 3 | 2.48 | 48 | 4.44 |
| Pākehā/European | 348 | 44.73 | 0 | 0.00 | <0.0001 | 99 | 81.82 | <0.0001 | 447 | 41.31 |
| Missing/no response | 13 | 1.68 | 0 | 0.00 | <0.0001 | 3 | 2.48 | 16 | 1.48 |
| School decile rating | | | | | | | | | <0.0001 |
| High | 545 | 70.05 | 80 | 43.72 | <0.0001 | 54 | 44.63 | <0.0001 | 679 | 62.75 |
| Medium | 159 | 20.44 | 54 | 29.51 | <0.0001 | 48 | 39.67 | 261 | 24.12 |
| Low | 24 | 3.08 | 38 | 20.77 | <0.0001 | 9 | 7.44 | 71 | 6.56 |
| Missing | 50 | 6.34 | 11 | 6.01 | <0.0001 | 10 | 8.26 | 71 | 6.56 |
| Attended school in Auckland | 565 | 72.62 | 115 | 62.84 | 0.0017 | 48 | 39.67 | <0.0001 | 728 | 67.28 |
| Completed bridging programme | 8 | 1.03 | 41 | 22.40 | <0.0001 | 1 | 0.83 | 1.0000 | 50 | 4.62 |
| Admitted as school leaver (Year 1) | | | | | | | | | <0.0001 |
| Graduate | 121 | 15.55 | 32 | 17.49 | <0.0001 | 36 | 29.75 | 189 | 17.47 |
| Undergraduate | 657 | 84.45 | 151 | 82.51 | <0.0001 | 85 | 70.25 | 893 | 82.53 |
| Graduated (yes) | 735 | 94.47 | 146 | 79.78 | <0.0001 | 111 | 91.74 | 0.2343 | 992 | 91.68 |
| Continuous variables | | | | | | | | | |
| Age at admission (Year 2) | 20.02 | 3.10 | 20.98 | 3.97 | 0.0014 | 21.07 | 3.80 | 0.0041 | 20.3 | 3.37 |
| Admission GPA/GPE | 8.12 | 0.99 | 6.04 | 1.29 | <0.0001 | 7.72 | 0.88 | 0.0002 | 7.72 | 1.29 |

Cohort 2002–2009 comprises students who matriculated into Year 2 of the MBCHB (or Bachelor of Human Biology BHB) programme within FMHS from 2002 to 2009 inclusive, excluding international entry students. Students who repeated Year 2 have been recorded in their first 2 years. All variables have been compared for the GENERAL versus MAPAS categories and for the General versus RRAS categories. Categorical variables have been tested using χ² test or Fisher’s exact test, where necessary. Continuous variables have been tested using the analysis of variance model, with adjustment for multiple comparisons.

GPA, Grade Point Average; GPE, Grade Point Equivalent; MAPAS, Māori and Pacific Admission Scheme; ROMPE, Rural Origin Medical Preferential Entry.
(30%, p<0.0001), admission into first year as an SL (59%, p<0.0001) and entry pathway into medicine (31% graduate, 69% undergraduate, p<0.0002). The average admission GPA/GPE was approximately half a point lower for ROMPE compared with GENERAL students (7.74, SD 1.19, p<0.0001).

Of the 1082 students who completed the programme in the study period (ie, admitted between 2002 and 2009), 95% (735/778) of GENERAL, 92% (111/121) of ROMPE and 80% (146/183) of MAPAS students graduated from MBChB. For the total cohort (admitted between 2002 and 2012), the mean Year 2–3 GPA was 0.35 (SD 1.52) for GENERAL, 5.82 (SD 1.65, p=0.0013) for ROMPE and 4.33 (SD 1.56, p<0.0001) for MAPAS students. Table 2 presents the multiple regression analysis findings for the total cohort.

Graduated from medicine
In the unadjusted model, MAPAS students had significantly lower odds of graduating from intended programme compared with GENERAL students (OR: 0.231, 95% CI: 0.144 to 0.371). This pattern remained after controlling for age, gender and year of admission, that is, model 1 (OR: 0.235, CI: 0.143 to 0.386). The odds of MAPAS students graduating in comparison to GENERAL students improved with the addition of medium and low school decile, that is, model 2 (OR: 0.291, CI: 0.165 to 0.513) and having attended a school out of Auckland or being admitted into first year as an SL, that is, model 3 (OR: 0.296, CI: 0.166 to 0.526, p=0.0002). The addition of having attended a bridging programme increased the odds of MAPAS students graduating from medicine by a further 14% in comparison to GENERAL students, that is, model 5 (OR: 0.440, CI: 0.231 to 0.841). When entry pathway into medicine as a graduate and a higher decile school, a school outside of Auckland and admission into first year as a mature student reduced the difference in GPA slightly, that is, model 3 (OR: 1.899, CI: −2.076 to −1.702). Having attended a bridging programme prior to medical study further reduced the difference in GPA between MAPAS and GENERAL students, that is, model 4 (OR: −1.724, CI: −1.914 to −1.533). When both graduate entry admission and admission GPA/GPE were added in model 5, no significant difference in Year 2–3 GPA was observed between the admission categories (OR: 0.103, CI: −0.103 to 0.309). These findings suggest that having attended a bridging programme, entering medicine as a graduate and a higher admission GPA/GPE are associated with improved performance for MAPAS compared with GENERAL students in the early years of the medical programme.

In the unadjusted model, the average difference between Year 2–3 GPA was approximately half a point lower for ROMPE compared with GENERAL students (OR: −0.449, CI: −0.668 to −0.23). This general pattern remains for models 1–4. When all predictor variables were taken into account in model 5, the mean difference in Year 2–3 GPA became non-significant (OR: −0.142, CI: −0.326 to 0.043). These findings suggest that admission as a graduate and admission GPA/GPE are the major contributors to the GPA difference between ROMPE and GENERAL students.

Table 3A and B presents the multiple regression analysis findings for the subcohort analyses for the MAPAS and GENERAL cohorts.

MAPAS subcohort
After controlling for predefined confounders (eg, gender, age, ethnicity, year of admission) and all significant predictors, that is, model 5, the odds of a MAPAS student graduating from medicine was 86% lower for those MAPAS students who attended a bridging programme versus those who did not (OR: 0.141, CI: 0.042 to 0.468, p=0.0014) and 83% lower for MAPAS students who entered medicine via the graduate pathway versus the undergraduate pathway (OR: 0.170, CI: 0.043 to 0.681, p=0.0123). The odds of graduating increased by 1.8 times for every point increase in admission GPA/GPE (OR: 1.758, CI: 1.05 to 2.944, p=0.0319). There were mixed findings for school decile across the models and this variable was not significant in the final model that included admission GPA/GPE and entry pathway.

For MAPAS students, the year 2–3 GPA was similar for students regardless of whether or not they had attended a bridging programme (−0.927, CI: −1.209 to 0.645, p<0.0001) and was 25% higher for every point increase in admission GPA/GPE (0.754, CI: 0.647 to 0.861, p<0.0001). School decile rating was not a significant predictor in the final model for the MAPAS cohort.
| Model | Predictor variable (ref) | Competition | Graduated (n=1082) (2002–2009 cohort) | Year 2–3 GPA (n=1676) (2002–2012 cohort) |
|-------|-------------------------|-------------|--------------------------------------|------------------------------------------|
|       |                         | Overall p-Value | OR | 95% CI          | Overall p-Value | Mean difference | 95% CI          |
| Unadj. | MAPAS                  | <0.0001      | 0.23 | 0.14 to 0.37 | <0.0001 | −1.93 | −2.11 to 1.76 |
|       | ROMPE                  | 0.65         | 0.32 to 1.33 | −0.45 | −0.67 to 0.23 |
| 1     | MAPAS                  | <0.0001      | 0.24 | 0.14 to 0.39 | <0.0001 | −1.99 | −2.17 to 1.82 |
|       | ROMPE                  | 0.70         | 0.33 to 1.50 | −0.54 | −0.76 to 0.32 |
|       | n=1011*                |              |      |                | n=1586* |              |                |
| 2     | MAPAS                  | 0.0001       | 0.29 | 0.16 to 0.51 | <0.0001 | −1.94 | −2.12 to 1.75 |
|       | ROMPE                  | 0.63         | 0.28 to 1.44 | −0.54 | −0.77 to 0.31 |
|       | School decile (high 8–10) Medium (4–7) | 0.0032 | 0.56 | 0.32 to 0.98 | 0.0279 | −0.16 | 0.00 to 0.31 |
|       | Low (1–3)              | 0.29         | 0.14 to 0.61 | −0.32 | −0.61 to 0.03 |
| 3     | MAPAS                  | 0.0002       | 0.30 | 0.17 to 0.53 | <0.0001 | −1.89 | −2.08 to 1.70 |
|       | ROMPE                  | 0.48         | 0.21 to 1.12 | −0.53 | −0.76 to 0.31 |
|       | School decile (high 8–10) Medium (4–7) | 0.0022 | 0.52 | 0.29 to 0.93 | 0.0454 | −0.15 | 0.00 to 0.31 |
|       | Low (1–3)              | 0.28         | 0.13 to 0.59 | −0.29 | −0.58 to 0.00 |
|       | Auckland school (yes)  No | 0.0030 | 2.67 | 1.40 to 5.09 | – | – | – |
|       | Type of admission AA   | 0.0430 | 0.53 | 0.29 to 0.10 | 0.0004 | −0.34 | −0.53 to 0.15 |
| 4     | MAPAS                  | 0.0182       | 0.44 | 0.23 to 0.84 | <0.0001 | −1.72 | −1.91 to 1.53 |
|       | ROMPE                  | 0.42         | 0.18 to 0.10 | −0.61 | −0.83 to 0.39 |
|       | School decile (high 8–10) Medium (4–7) | 0.0096 | 0.58 | 0.32 to 1.05 | – | – | – |
|       | Low (1–3)              | 0.31         | 0.14 to 0.68 | − | – | – |
|       | Auckland school (yes)  No | 0.0062 | 2.52 | 1.30 to 4.88 | – | – | – |
|       | Type of admission AA   – | – | – | – | – | – | – |
|       | Bridging Programme (no) Yes | <0.0001 | 0.16 | 0.07 to 0.36 | <0.0001 | −1.24 | −1.56 to 0.91 |
| 5     | MAPAS                  | 0.1251       | 1.68 | 0.74 to 3.83 | 0.1306 | 0.10 | −0.10 to 0.31 |
|       | ROMPE                  | 0.56         | 0.23 to 1.37 | −0.14 | −0.33 to 0.04 |
|       | School decile (high 8–10) Medium (4–7) | 0.0276 | 0.66 | 0.35 to 1.23 | – | – | – |
|       | Low (1–3)              | 0.31         | 0.13 to 0.74 | – | – | – |
After controlling for predefined confounders (e.g., gender, age, ethnicity, year of admission) and all significant predictors, the odds of a GENERAL student graduating from medicine was lower for students who attended a low decile (OR: 0.137, CI: 0.031 to 0.6, p=0.0098) or medium decile school (OR: 0.384, CI: 0.164 to 0.898).

**Table 2** Continued

| Model | Predictor variable (ref) | Comparison | Graduated (n=1082) (2002–2009 cohort) | Year 2–3 GPA (n=1676) (2002–2012 cohort) |
|-------|--------------------------|------------|--------------------------------------|----------------------------------------|
|       |                          | Overall p-Value | OR | 95% CI | Overall p-Value | Mean difference | 95% CI   |
|       |                          |                |    |        |                |                |          |
| 1     | Ethnicity (reference group Māori for MAPAS, Pākehā for GENERAL) | Māori – – – | 0.0030 | 2.88 | 1.43 to 5.79 | – – – | – – – |
|       |                          | Pacific       | 0.0035 | 0.285 | 0.123 to 0.662 | – – – | – – – |
|       |                          | Asian – – –   |        |        |                |        |          |
|       |                          | Other/missing | – – | – – | – – – | – – – | – – – |
| 5     | School decile (high 8–10) | Medium (4–7) | 0.0098 | 0.384 | 0.164 to 0.898 | – – – | – – – |
|       |                          | Low (1–3)     | 0.137 | 0.031 | 0.031 to 0.600 | – – – | – – – |
|       | Auckland school (yes)    | No            | 0.0127 | 4.571 | 1.256 to 16.629 | – – | – – |
|       | Type of admission (SL)   | AA            | – – | – – | – – – | – – – | – – – |
|       | Bridging programme (no)  | Yes           | 0.0014 | 0.141 | 0.042 to 0.468 | – – | – – |
|       | Entry pathway (undergraduate) | Graduate | 0.0123 | 0.170 | 0.043 to 0.681 | – | – |
|       | Admission GPA/GPE        | Per point increase | 0.0319 | 1.758 | 1.050 to 2.994 | <0.0001 | 2.020 | 1.460 to 2.796 |

**Table 3A** Logistic regression results for Graduated (2002–2009 cohort) academic outcome for MAPAS and GENERAL subgroups

| Model | Predictor variable (ref) | Comparison | Graduated (2002–2009 cohort) | MAPAS (n=181) | Graduated (2002–2009 cohort) | GENERAL (n=778) |
|-------|--------------------------|------------|-----------------------------|---------------|-----------------------------|-----------------|
|       |                          | Overall p-Value | OR | 95% CI | Overall p-Value | OR | 95% CI |
|       |                          |                |    |        |                |    |        |
| 1     | Ethnicity (reference group Māori for MAPAS, Pākehā for GENERAL) | Māori – – – | 0.0010 | 0.175 | 0.047 to 0.648 | – – – | – – – |
|       |                          | Pacific       | 0.0035 | 0.285 | 0.123 to 0.662 | – – – | – – – |
|       |                          | Asian – – –   |        |        |                |        |          |
|       |                          | Other/missing | – – | – – | – – – | – – – | – – – |
| 5     | School decile (high 8–10) | Medium (4–7) | 0.0098 | 0.384 | 0.164 to 0.898 | – – – | – – – |
|       |                          | Low (1–3)     | 0.137 | 0.031 | 0.031 to 0.600 | – – – | – – – |
|       | Auckland school (yes)    | No            | 0.0127 | 4.571 | 1.256 to 16.629 | – | – |
|       | Type of admission (SL)   | AA            | – – | – | – – – | – – – | – – – |
|       | Bridging programme (no)  | Yes           | 0.0014 | 0.141 | 0.042 to 0.468 | – | – |
|       | Entry pathway (undergraduate) | Graduate | 0.0123 | 0.170 | 0.043 to 0.681 | – | – |
|       | Admission GPA/GPE        | Per point increase | 0.0319 | 1.758 | 1.050 to 2.994 | <0.0001 | 2.020 | 1.460 to 2.796 |
decile school (OR: 0.384, CI: 0.164 to 0.898, p=0.0098) compared with high school decile. Increasing admission GPA/GPE was strongly associated with increased odds of graduating (OR: 2.020, CI: 1.46 to 2.796, p<0.0001). The Year 2–3 GPA was lower for graduate entry GENERAL students compared with undergraduate entry (0.577, CI: 0.377 to 0.777, p=0.0036) with similar outcomes observed for bridging programme attendance (−1.083, CI: −1.812 to −0.355, p=0.0036) and admission GPA/GPE (0.977, CI: 0.891 to 1.063, p=<0.0001). School decile rating was not a significant predictor of early academic outcomes for the GENERAL cohort.

**DISCUSSION**

This study, based on 1676 medical students over a 10-year period compared outcomes and predictor variables of those admitted via two equity-admission pathways with those in the general admission pathway. To our knowledge, this is the first report in the literature describing programme level outcomes to this detail. The descriptive data confirm that it is possible to admit significant numbers of students via these pathways and have most successfully complete the programme. Nearly all students with Māori and Pacific ethnicity entered via the MAPAS pathway. Furthermore, the MAPAS and ROMPE pathways each contained higher proportions of students from lower socioeconomic backgrounds and students who attended schools out of Auckland. These findings underscore the importance of having equity pathways or targets, as it unlikely many of the MAPAS students, and some of the ROMPE students would have been successful in the highly competitive selection process for GENERAL students. Furthermore, to provide workforce benefit, students need to complete the programme. Encouragingly, despite marked differences in background and prior performance there was only a 12%–15% difference in the proportion of MAPAS students who graduated in the study period compared with ROMPE or GENERAL admission students respectively. Our hypotheses that markers of socioeconomic status, transitioning factors, bridging programme attendance and academic preparation are likely to impact on both short-term and long-term academic outcomes were confirmed, although findings are mixed within and across the entry pathways. When looking within the MAPAS cohort, the odds of a MAPAS student graduating (compared with another MAPAS student) improved with non-bridging programme attendance and most likely reflect cohort differences in admission GPA/GPE. In contrast, our findings suggest that having attended a bridging programme, entering medicine as an undergraduate and higher admission GPA/GPE are the major contributors to reducing the

| Table 3B  | Linear regression results for Year 2–3 GPA (2002–2012 cohort) academic outcome for MAPAS and GENERAL subgroups |
|-----------|-------------------------------------------------------------------------------------------------------------|
| **Model** | **Predictor variable (ref)**                                                                                   | **Comparison**                                      | **Year 2–3 GPA (2002–2012 cohort)** | **Year 2–3 GPA (2002–2012 cohort)** |
|           |                                                                                                              |                                                                                                         |  **MAPAS (n=315)** |  **GENERAL (n=1167)** |
| 1         | Ethnicity                                                                                                     | Māori                                                | **Overall** | <0.0001 | **Mean difference** | <0.0001 | **95% CI** | <0.0001 | **Mean difference** | <0.0001 | **95% CI** |
|           | (reference group                                                                                              | Pacific                                              | 0.0109 | −0.425 | −1.018 to 0.168 | −0.616 | −1.201 to 0.031 |
|           | Māori for MAPAS, Pākehā/ European for GENERAL                                                                  | Asian                                                | −        | −       | −                  | −2.226 | −2.939 to 1.513 |
|           |                                                                                                              | Other/missing                                        | −        | −       | −                  | −0.191 | −0.358 to 0.025 |
|           |                                                                                                              |                                                                                                         | −        | −       | −                  | −0.524 | −0.852 to 0.196 |
| 5         | School decile (high 8–10)                                                                                     | Medium (4–7)                                         | n=302    |        |        | n=1104    |        |        |
|           | Low (1–3)                                                                                                     |                                                                                                         | −        | −       | −                  | −        | −        |
|           | Auckland school (yes)                                                                                          | No                                                    | −        | −       | −                  | −        | −        |
|           | Type of admission (SL)                                                                                        | AA                                                    | −        | −       | −                  | −        | −        |
|           | Bridging programme (no)                                                                                       | Yes                                                   | <0.0001 | −0.927 | −1.209 to −0.654 | 0.0036 | −1.812 to 0.355 |
|           | Entry pathway (undergraduate)                                                                                 | Graduate                                              | −        | −       | −                  | <0.0001 | 0.577    | 0.377 to 0.777 |
|           | Admission GPA/GPE Per point increase                                                                           |                                                        | <0.0001 | 0.754  | 0.647 to 0.861    | <0.0001 | 0.891    | 1.063 |

Logistic regression model has controlled for year of admission, gender and age at admission. Predefined predictors were added to the baseline model in sequential order to estimate their joint independent effects on the outcome. Each model was initially run with all the specified predictors, then rerun with stepwise selection to include significant predictors only and obtain final estimates of effect size. Model-adjusted estimates of ORs (compared with the reference level), 95% CI and associated individual p-values (in symbols) were reported. AA, alternative admission (ie, non-school leaver); GPA, Grade Point Average; GPE, Grade Point Equivalent; MAPAS, Māori and Pacific Admission Scheme; SL, school leaver.
GPA difference observed between MAPAS and GENERAL students in the early, non-clinical phase of medical training.

This study represents a comprehensive analysis of academic outcomes for equity admission pathways into medicine within NZ. Similarly, this study explores academic outcomes for one of the largest cohorts of indigenous medical students available internationally. We acknowledge that this study was confined to a single medical programme and that the results may not be generalisable to other programmes or tertiary institutions. In particular, the comprehensive nature of the MAPAS programme with respect to student admission and retention support may not be reflected in other tertiary contexts. Like similar measures elsewhere (eg, participation of local areas (P FAR) classification in England), the use of secondary school decile as a proxy for socioeconomic position relies on an area-level indicator of deprivation and may not directly reflect the socioeconomic position of each individual student or their family. Despite this, other school factors (eg, student attainment, aspirations for future study) have been linked to school decile suggesting that individual students will have been exposed to direct school effects. This study did not explore the effect of medical interview outcomes due to different processes of selection being used across equity and general admission pathways. The study period spans across and before periods of significant change within the MAPAS and ROMPE pathways with respect to admissions processes (ie, selection methods and eligibility). Therefore, study findings should be interpreted cautiously as ‘historical’ markers of equity programme delivery or performance rather than accurate representations of the equity processes in operation today.

Our findings are consistent with the existing literature base that GPA at the point of admission is the strongest predictor of academic outcomes within the medical programme. In a critical appraisal of studies examining medical school failure, O’Neill and colleagues found that lower entry qualifications at admission were linked to higher failure rates. However, they note that many studies did not control for confounding factors, were mostly focused on student attributes, with few studies examining the role of the institution. The fact that 80% of MAPAS students completed medicine despite being admitted with an average GPA approximately two points lower than other medical students is encouraging. This suggests that while GPA at admission is important, other unmeasured factors may be contributing to our findings. Student pastoral and financial issues (likely to be significant for indigenous students given their socioeconomic and demographic profile), psychological characteristics, student learning styles and relevant medical curricula or structural factors may also play a role. As noted by Mathers and Parry, graduate applicants to medicine have complex needs arising from their personal social, family and economic circumstances that may affect their academic performance. The University of Auckland’s commitment to respond to these student factors (via the provision of comprehensive admission, pastoral and academic support) may be contributing to our outcomes observed, particularly for graduate entry and bridging programme students admitted under MAPAS. The positive effect of bridging programme exposure has also been noted elsewhere. However within the MAPAS cohort, those students who did not require additional academic support via a bridging programme experienced better academic outcomes. Therefore, our findings reinforce the need for ongoing bridging programme delivery alongside the elimination of educational inequities for Māori and Pacific students (for more information please see https://www.fmhs.auckland.ac.nz/en/faculty/for/future-undergraduates/undergraduate-study-options/certshc.html). The association between secondary school decile rating (a marker of socioeconomic status and school characteristics) and academic outcomes had mixed results and are unlikely to explain the differences observed by admission pathway. Although school decile has been linked to first year academic outcomes for Māori, our findings may reflect the fact that school characteristics have been noted to have less impact on student achievement at the higher end of the achievement scale, that is, GPA ≥ 4. However, the strong association between lower school decile and reduced odds of graduating for the GENERAL admission students challenges this conclusion, differs from other research and is of concern.

Our study reinforces the existing evidence that equity-targeted admission programmes, inclusive of variations in academic threshold for entry, can support a widening participation agenda within medicine. However, tertiary institutions and society at large must accept that ethnic inequities in educational outcomes and rural workforce development needs should be accounted for within admission pathways and retention support. Providing comprehensive academic and pastoral assistance to equity admission and lower socioeconomic students who are operating within complex and academically demanding contexts remains paramount. While differences in academic thresholds for equity groups appears necessary, it is often criticised as being ‘politically correct’, providing ‘preferential treatment’ to one group or individual over another and has not been universally welcomed by the public or the profession. Bacchi notes that the framing of widening participation as ‘preferential treatment’ ‘undermines the legitimacy of the reform and reduces its impact, limiting the kinds of reforms ‘permitted’ and alienating those who are targeted. This undoubtedly serves the interests of those who profit under current social arrangements’ (p144).

Given this context, it is perhaps not surprising that while medical schools strive to increase diversity and meet the goals of a widening participation agenda, successful implementation is influenced by contextual factors associated with institutional leadership, resource allocation and external stakeholder pressure. Razack et al note that
while the development of social accountability policy has occurred, medical schools appear to be challenged by the implementation of these policies within student recruitment and selection processes.6 The study responds to calls for open and inclusive discussions in order to advance admissions practice aiming to enhance social justice and widening participation agendas.6

Additional research is warranted (eg, inclusion of secondary school outcomes, non-cognitive testing and medical interview data beyond 2012). Similarly, exploring the effect of institutional attributes should also be considered.1,7 Evidence suggests that tertiary and medical school environments may have different effects on indigenous and ethnic minority students who have reported that their ethnicity adversely affects their medical school experience,62 have described experiences of racism from peers and clinical educators,28 and are adversely affected by an ‘othering’ medical curriculum that either stereotypes indigenous culture and society or fails to reflect indigenous realities altogether.35, 64–66 Exploring the impact of these variables on differential academic outcomes for equity admission pathways may require qualitative methods to complement additional quantitative analyses. The impact of a widening participation agenda within medicine must also begin to look beyond the number of students admitted and graduated and extend the analysis to post-graduate clinical contexts including the effect of a diverse health workforce on patient and community outcomes.1 The ultimate aim of equity-targeted admissions pathways into medicine is to enhance healthcare delivery, improve health outcomes and eliminate inequities for underserved communities. Understanding when and how this can be achieved remains a challenge for many countries worldwide.

Author affiliations
1Te Kupenga Hauora Māori, Faculty of Medical and Health Sciences, University of Auckland, Auckland, New Zealand
2Department of Statistics, Faculty of Science, University of Auckland, Auckland, New Zealand
3Department of Medicine, Faculty of Medical and Health Sciences, University of Auckland, Auckland, New Zealand
4Faculty of Medical and Health Sciences, University of Auckland, Auckland, New Zealand
5Medical Programme Directorate, Faculty of Medical and Health Sciences, University of Auckland, Auckland, New Zealand
6Medical interview data beyond 2012). Similarly, exploring the effect of institutional attributes should also be considered.1,7 Evidence suggests that tertiary and medical school environments may have different effects on indigenous and ethnic minority students who have reported that their ethnicity adversely affects their medical school experience,62 have described experiences of racism from peers and clinical educators,28 and are adversely affected by an ‘othering’ medical curriculum that either stereotypes indigenous culture and society or fails to reflect indigenous realities altogether.35, 64–66 Exploring the impact of these variables on differential academic outcomes for equity admission pathways may require qualitative methods to complement additional quantitative analyses. The impact of a widening participation agenda within medicine must also begin to look beyond the number of students admitted and graduated and extend the analysis to post-graduate clinical contexts including the effect of a diverse health workforce on patient and community outcomes.1 The ultimate aim of equity-targeted admissions pathways into medicine is to enhance healthcare delivery, improve health outcomes and eliminate inequities for underserved communities. Understanding when and how this can be achieved remains a challenge for many countries worldwide.

Contributors
EC (senior lecturer medical) is the guarantor of the study; led the study design, drafting and revising of the manuscript. PP contributed to the study design and provided senior statistical expertise for data analysis; contributed to drafting and revising the manuscript. PR (Tumaki, deputy dean Māori) provided senior Māori educational, institutional and Kaupapa Māori expertise and contributed to the study design, drafting and revising the manuscript. All authors read and approved the final manuscript for submission.

Funding
EC was supported by Te Kete Hauora, Ministry of Health (New Zealand) to conduct this research via the provision of a Research Fellowship (contract 414953/337535/00).

Competing interests
None declared.

Ethics approval
The study was approved by the UoA Human Participants Ethics Committee (UAHPEC) (reference 8110).

Provenance and peer review
Not commissioned; externally peer reviewed.

Data sharing statement
No additional data are available.

Open Access
This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/

© Article author(s) (or their employer(s) unless otherwise stated in the text of the article) 2017. All rights reserved. No commercial use is permitted unless otherwise expressly granted.

REFERENCES
1. Baxter C, Baxter D, Baxter M. Widening participation in medicine: moving beyond the numbers. Med Educ 2015;49:7–20.
2. Powis D, Hamilton J, McManus IC. Widening access by changing the criteria for selecting medical students. Teach Teach Educ 2007;23:1225–45.
3. Cleland JA, Nicholson S, Kelly N, et al. Taking context seriously: explaining widening access policy enactments in UK medical schools. Med Educ 2015;49:25–35.
4. Garlick PB, Brown G. Widening participation in medicine. BMJ 2008;336:1111–3.
5. Cooper LA, Beach MC, Johnson RL, et al. Delving below the surface. Understanding how race and ethnicity influence relationships in health care. J Gen Intern Med 2006;21 (Suppl. 1):S21–S27.
6. van Ryn M, Fu SS. Paved with good intentions: do public health and human service providers contribute to racial/ethnic disparities in health? Am J Public Health 2003;93:248–55.
7. Betancourt JR, Green AR, Carrillo JE, Emilio Carrillo J, et al. Cultural competence and health care disparities: key perspectives and trends. Health Aff 2005;24:499–505.
8. Cantor JC, Miles EL, Baker LC, et al. Physician service to the underserved: implications for affirmative action in medical education. Inquiry 1996;33:167–80.
9. Larkins S, Michielsen K, Iputo J, et al. Impact of selection strategies on representation of underserved populations and intention to practise: international findings. Med Educ 2015;49:60–72.
10. Whita DK, Orfield G, Silen W, et al. Educational benefits of diversity in medical school: a survey of students. Acad Med 2003;78:460–6.
11. Hung R, McClenon J, Henderson A, et al. Student perspectives on diversity and the cultural climate at a U.S. medical school. Acad Med 2007;82:184–92.
12. Elam CL, Johnson MM, Wiggs JS, et al. Diversity in medical school: perceptions of first-year students at four southeastern U.S. medical schools. Acad Med 2001;76:80–5.
13. van Ryn M, Hardeman R, Phelan SM, et al. Medical School Experiences Associated with Change in Implicit Racial Bias among 3547 students: a Medical Student CHANGES Study Report. J Gen Intern Med 2015;30:1748–56.
14. Seyan K, Greenhalgh T, Doling D. The standardised admission ratio for measuring widening participation in medical schools: analysis of UK medical school admissions by ethnicity, socioeconomic status, and sex. BMJ 2004;328:1545–6.
15. Ready T. The Impact of Affirmative Action on Medical Education and the Nation’s Health. In: Orfield G, Kurlander M, eds. Diversity challenges evidence on the impact of Affirmative Action. Cambridge: Harvard Educational Review, 2001:205–19.
