| | | MEDICARE CARD NUMBER | | | |
|---|--|----------------------|---|------------------------------|--------------|
| COMPASS Future directions In cervical screening VCS Pathology | | PA | | PATHO | LOGY REQUEST |
| Patient Surname Given Na | Given Names | | Date of Birth | Date of Birth Your Reference | |
| Patient Address | | Previous Surname | | | |
| LAB COPY | | | | | |
| Tests Requested s.D. Clinical Notes Urgent Phone Fax By Time: | Is the patient of Aboriginal or Torres Strait Islander origin? Aboriginal Torres Strait Islander Aboriginal and Torres Strait Isla Not Aboriginal or Torres Strait Isla Not Aboriginal or Torres Strait Isla In which country was the patient Does the patient speak a language ott English at home? (If more than one lan indicate the one that is spoken most oft No, English only Yes, othe [please specify] Practitioner' | | Pregnant/Post partum Hysterectomy HRT IUCD Specimen site Same day colposcopy | No Yes | |
| Phone/Fax No Private Schedule Bulk Bill (Complete Medicare Assignment) Vet Affairs No | | | | | |
| COPY REPORTS TO: | | Requesting Practit | ioner (Provider number, Su | rname, Initials and | Address) |
| MEDICARE ASSIGNMENT (Section 20A of the Health Insurance Act 1973). I assign my right to benefits to the approved pathology practitioner who will render the requested pathology service(s) practitioner only (please tick). Reason patient unable to sign. | | Patient's Signatur | e | | Date |
| Compass Trial: Study Participant Consent Form - Participation involves the following: Researchers are allowed to access my relevant medical records • I am aged at least 25 and born on or after 1st July 1980 • My participation in the trial is entirely voluntary and I can withdraw at any stage. This will not affect my relationship with my health practitioner, VCS Pathology or Cancer Council NSW • Any information relating to my participation in the study is strictly confidential. I agree that the results can be published, provided I cannot be identified • Any residual samples will be stored and may be used for the development of new HPV testing technologies. Ethical approval will be obtained prior to undertaking any further research studies using these samples. I understand that I will not have access to these test results • I have any questions I can contact the Compass trial hotline on 1800 611 635 or visit www.compasstrial.org.au • If I have any concerns or complaints about the study, I can contact the Research Governance Officer, Bellberry Limited on 08 8361 3222 • I may be contacted in the future and asked to complete a questionnaire or receive a study newsletter • I understand that the Compass Register will undertake the usual functions of my local Pap test register • I consent to the Compass investigators opting me out of receiving correspondence from my local Pap test register; on my behalf • I understand that I will be followed on the Compass register for the duration of the triat. I agree to participate in the Compass Trial and understand that I will be followed for 5 years. • I understand I will be randomly allocated to having my cervical screening sample tested by one of two methods; cytology [the test used up until December 2017) in the cervical screening program] or HPV DNA testing [the new test used from December 2017] I d o not wish for my left-over cervical sample to be used for any further research or for any HPV test validation purposes. PO Box 178 Cartton South VIC 3053 P: 03 9 | | | | | |
| [name] <i>please print clearly</i> | | | | | |
| Name | Phone | | Signature | | Date |
| Email | Mobile | | Х | | |

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