

 MISSOURI DEPARTMENT OF HEALTH AND SENIOR SERVICES SHOW ME HEALTHY WOMEN (SMHW) SCREENING REPORT		P. O. Box 57 Jefferson City, MO 65102-057 (573) 522-284	
ENROLLMENT SITE/SATELLITE SITE (NAME AND ADDRESS)		REFERRING PROVIDER (FOR DIRECT BILLING)	
A. PERSONAL DATA			
NAME (LAST, FIRST, MIDDLE INITIAL)		SOCIAL SECURITY NUMBER	
DATE OF BIRTH MM / DD / YYYY	CLIENT ELIGIBILITY VERIFIED ☐ Yes ☐ No	INSURANCE COVERAGE ☐ Yes ☐ No	DEDUCTIBLE MET ☐ Yes ☐ No
		REFERRAL FEE ☐	MEDICARE ☐ Part A ☐ Part A and B
VISIT TYPE ☐ Initial ☐ Annual ☐ Initial CBE only ☐ Annual CBE only ☐ Mammogram only		Height ft. in.	Weight lbs.
		Blood Pressure 1st Reading / / 2nd Reading / / Average / /	
B. BREAST CANCER SCREENING			
☐ Reporting Only			
B 1. Does client report any BSE symptoms? ☐ Yes ☐ No (If "YES" complete B2.)			
B 2. Symptoms Reported By Client (Check any that apply. If 1, 2, 3 or 4B is checked, may have two (2) diagnostics at clinician's discretion.)			
☐ (1) Lump ☐ (2) Nipple discharge ☐ (3) Skin changes (dimpling, retraction, new nipple inversion, ulceration, Paget's disease) ☐ (4A) Pain/Tenderness - 1st occurrence ☐ (4B) Pain/Tenderness - 2nd occurrence ☐ (5) Other (specify) _____			
B 3. CBE within normal limits and findings Present at CBE (check yes or no and one explanation) Date of CBE / / (MM/DD/YYYY)			
☐ Yes ☐ Within normal limits ☐ (1) Benign finding (fibrocystic changes, diffuse lumpiness, clearly defined thickening, tenderness or nodularity) ☐ No - Suspicious for cancer (Any checked findings requires completion of two (2) diagnostic procedures entered on purple breast form.) ☐ (2) Discrete palpable mass (includes masses that may be diffuse, poorly defined thickening, cystic or solid) ☐ (3) Nipple discharge ☐ (4) Nipple or areolar scaliness or erythema ☐ (5) Skin dimpling retraction; new nipple inversion; peau d'orange; ulceration; one breast lower than usual; prominent veins, unilateral; unusual increase in size, unilateral ☐ (6) Enlarged, tender, fixed or hard palpable supraclavicular, infraclavicular or axillary lymph nodes; also swelling of upper arm ☐ Focal pain and tenderness			
Rescreen CBE Planned ☐ Yes ☐ No / / (must be less than 10 months) MM YYYY			
Diagnostic Workup Planned ☐ Yes ☐ No / / (must be less than 10 months) MM YYYY			
B 4. High Risk for Breast Cancer ☐ (1) Yes* ☐ (2) No ☐ (9) Not assessed/Unknown * At least one must be met: BRCA Mutation, First Degree Relative BRCA Carrier, or Greater Than 20-26 Percent Lifetime Risk			
B 5. Mammogram			
Previous mammogram ☐ Yes ☐ No ☐ Unknown Date of last mammogram / / MM YYYY Date of this mammogram / / MM DD YYYY			
Type of mammogram ☐ Screening ☐ Diagnostic ☐ Tomosynthesis Method used for mammogram ☐ Digital ☐ Conventional			
Mammography provider facility (facility name / city) ☐ Mammogram Van			
☐ (4) Mammogram not done or CBE done and diagnostic workup planned ☐ (1) Routine screening mammogram ☐ (2) Mammogram performed to evaluate symptoms: ☐ Personal history of breast cancer ☐ Previous abnormal mammogram results (rescreen) ☐ (5) Cervical record only, no breast service provided ☐ (6) Referred to direct biller ☐ (3) Abnormal mammogram done by a non-program funded provider, patient referred in for diagnostic evaluation (Enter results in Mammogram field as Reporting Only) Date client referred for diagnosis. / / MM DD YYYY			
SMHW mammogram result (check one) (results with * require additional follow-up) ☐ Reporting Only			
Left Right (Indicate why only one breast had mammogram in COMMENTS) Normal ☐ (1) Negative (Category 1) ☐ (2) Benign Finding (Category 2) Left Right Abnormal ☐ (3) Probably Benign (Category 3) ☐ (4) Suspicious Abnormality (Category 4)* ☐ (5) Highly Suggestive of Malignancy (Category 5)* ☐ (7) Unsatisfactory-not interpreted, repeat (Not Paid) ☐ (14) Need evaluation or film comparison (Category 0)			
Further diagnostic planned for: (3) Probably Benign: ☐ Yes ☐ No			
Rescreen mammogram planned ☐ Yes ☐ No (must be less than 10 months) / / MM YYYY		Diagnostic work-up planned ☐ Yes ☐ No (must be less than 60 days) / / MM DD YYYY	
Referred for diagnostic testing/direct bill (physician / facility name) / / MM DD YYYY			
☐ MRI (High Risk ONLY. Prior authorization required.) MRI Type: ☐ Unilateral ☐ Not Done ☐ Bilateral		Report results here →	
MRI Results L R ☐ (1) Negative (Category 1) ☐ (2) Benign Finding (Category 2) ☐ (3) Probably Benign (Category 3) ☐ (4) Suspicious (Category 4) ☐ (5) Highly Suspicious (Category 5) ☐ (6) Known Malignancy (Category 6) ☐ (7) Incomplete (Category 0) ☐ (8) Results Pending ☐ (9) Not Done			

C. CERVICAL CANCER SCREENING (Indications for Pap Test)

☐ (6) Breast and Pelvic exam only (No Cervical Service)	☐ (1) Routine Pap test (<i>screening</i>) ☐ (2) Patient under surveillance for previous abnormal (<i>rescreen</i>) ☐ (5) Pap test not done. Patient proceeded directly for diagnostic work-up or HPV testing ☐ (4) Pap after primary HPV positive (+) ☐ (3) Non-program Pap referred in for diagnostic evaluation ☐ (9) Unknown	High Risk for Cervical Cancer ☐ (1) Yes ☐ (2) No ☐ (9) Not assessed/ unknown
		MM / DD / YYYY

C 1. Pelvic Exam ResultsPelvic Exam WNL? ☐ Yes ☐ No
(Additional information required in "No" selected, See C 2.)Hysterectomy? ☐ Yes ☐ No
☐ Cervix absent
☐ Cervix absent due to cervical cancer
(needs annual Pap test)
☐ Cervix present
☐ Reason for hysterectomy unknown

Date of Pelvic Exam MM / DD / YYYY

Reproductive Status (check one)

- ☐
- a) Premenopausal
-
- ☐
- b) Postmenopausal

C 2. Pelvic Exam Findings☐ Reporting Only

Findings Present at Pelvic Exam (check only one)

☐ 1) Cervix

- | | |
|---|---|
| ☐ a) Polyp | ☐ f) Ectropion |
| ☐ b) Leukoplakia (white lesions) | ☐ g) Stenotic OS |
| ☐ c) Friable | ☐ h) Cervical mass |
| ☐ d) Ulceration | ☐ i) Other: _____ |
| ☐ e) Exophytic growth | |

☐ 2) Exam Complicated by ObesityRescreen planned ☐ Yes ☐ No MM / YYYYDiagnostic planned ☐ Yes ☐ No MM / DD / YYYY
(must be less than 60 days)**C 3. Pap Test Results**☐ Reporting OnlyPrevious Pap test ☐ Yes ☐ No ☐ Unknown

Date of last Pap test MM / YYYY

Date of this Pap test MM / DD / YYYY

Specimen adequacy ☐ Satisfactory
☐ Unsatisfactory due to _____
☐ UnknownSpecimen type ☐ Conventional Smear
☐ Liquid Based☐ Annual Pap due to previous treatment for cervical cancer

Pap test result (check one) (Results with (*) require additional follow-up)

Normal ☐ (1) Negative for intraepithelial lesion or malignancy
☐ (2) Inflammatory/Infection/Reactive ChangesAbnormal ☐ (3) Atypical Squamous Cells of Undetermined Significance
(ASC-US)(May have HPV test)
☐ (4) Lowgrade SIL (HPV/Mild Dysplasia/CIN I)*
☐ (5) Atypical Squamous Cells, cannot exclude HSIL (ASC-H)*
☐ (6) Highgrade SIL (with features suspicious for invasion/CIN II-III/CIS)*

- ☐
- (7) Squamous Cell Cancer*
-
- ☐
- (8) Atypical Glandular Cells* (including atypical endocervical
-
- adenocarcinoma in situ and adenocarcinoma)
-
- ☐
- (9) Adenocarcinoma in situ*
-
- ☐
- (10) Adenocarcinoma*
-
- ☐
- (11) Other _____

Endocervical Cells ☐ Yes ☐ No**C 4. HPV Test Date** MM/DD/YYYY☐ Reporting OnlyIndication for HPV Test ☐ (1) Cotesting/Screening
☐ (2) Reflex
☐ (3) Not Done
☐ (9) Unknown

HPV Test Result

- | | |
|---|---|
| ☐ (1) Positive with genotyping
not done/unknown | ☐ (5) Positive with
negative genotyping |
| ☐ (2) Negative | ☐ (9) Unknown |
| ☐ (4) Positive with positive
genotyping | |

HPV DNA Genotype 16 or 18 Positive (Only report if PAP
negative and HPV High Risk Group Positive)

- ☐
- Yes
-
- ☐
- No
-
- ☐
- No Test Performed

Rescreen Pap planned ☐ Yes ☐ No MM / YYYY
(less than 10 months)Diagnostic work-up planned ☐ Yes ☐ No MM / DD / YYYY
(must be less than 60 days)Referred for diagnostic work-up/direct biller
(physician/facility name)

Date of next routine Pap screening MM / YYYY