Additional Risks
for Home Use of Pills
Prescribed by Telemedicine,
Delivered by Mail

Inadequate Screening
While women visiting a clinic can have a physical examination and an ultrasound, health care providers doing only an online or telephone interview rely on a woman giving them accurate, honest answers to questions about her last menstrual period, symptoms of ectopic pregnancy, allergies or other disqualifying conditions. If a woman is unaware of these contraindications, mistakes spotting for her last menstrual period, or does not yet recognize signs of ectopic pregnancy, these important factors may be missed and the abortion pills either may not work or may prove dangerous for her.1

Testing for Rh Factor
Failure to identify and treat Rh factor could mean the loss of future pregnancies.2

Higher Failure, Complication Rates
Less careful, less scientific screening will mean more women past the FDA recommended cutoff date of ten weeks, and thus a greater likelihood of complications or a failed or incomplete abortion.3

Less Assurance of Access to Emergency Care
Women doing screening by telemedicine and having pills shipped to remote locations may not necessarily have ready access to specialized surgical care from their prescriber or a nearby emergency care facility if they suffer sudden serious bleeding episodes or a ruptured ectopic pregnancy.4

Quality of Internet Medication
Women buying mifepristone and misoprostol from a foreign based internet pharmacy have no real assurance of product purity, dose, or efficacy and often receive little or no instructions on appropriate use.5

Sale to Dishonest, Deceptive Buyers
Online, telemedical, or telephonic screening and prescription allows those ordering to misrepresent their intentions or identities, potentially allowing these to be resold or given unknowingly to underage teens, unwilling or unscreened women with later pregnancies or conditions for whom these would not work or might prove dangerous.6

Difficulty knowing when you’re done.
Without a professional exam, it is possible a woman might bleed and cramp and think her abortion is completed, yet be mistaken.7

MIFEPRISTONE
SAFETY & EFFICACY
Quick Facts about the Abortion Pill

Drug-induced abortions are painful and bloody.
These pills work by shutting down the child’s life support system, initiating bleeding, and then stimulating powerful, painful contractions to expel the child and other contents from the uterus. Pain and bleeding are unavoidable parts of the process.8

Women lose more blood from a chemical than a surgical abortion.9

Chemical abortions take longer to complete than surgical ones.
Not counting recovery time, surgical abortions may take maybe 10 minutes to complete, so that a woman can be in and out of a clinic in a couple of hours10. Chemical abortions involve multiple drugs taken over a number of days and may take days or weeks to be fully completed.11

“Medication” abortions have a significant failure rate.
The FDA warns these drugs fail to deliver a complete abortion 2-7% of the time12.

The risk of failure and complications increases with gestational age.
The FDA originally limited use of mifepristone to women no more than 49 days after their last menstrual period (LMP) because of reduced efficacy and increased complications after that point.13 Years of field experience have confirmed this.14

Women see their aborted children’s bodies.
Identification of the embryo or fetus is one of the ways a woman is able to confirm the abortion pill has done its job, but it can also prove traumatic when women report seeing their child’s eyes, fists, or other body parts.15

There have been at least 28 deaths and thousands of injuries among American mifepristone patients.
The FDA reports that more than two dozen mifepristone patients in the U.S. have died after taking mifepristone and that thousands of others have suffered from complications such as hemorrhages, infections, and the rupture of previously undiscovered ectopic pregnancies, many requiring hospitalization and surgery.16

The warning signs of ectopic pregnancy are disturbingly similar to chemical abortion side effects.
Because women having drug-induced abortions normally face considerable abdominal pain and bleeding, patients and even doctors who have not seen an ultrasound have missed these danger signs of a rupturing ectopic pregnancy.17

Several mifepristone users came down with serious bacterial infections.
A number of women taking mifepristone died after contracting Clostridium sordellii, an anaerobic bacteria that thrives in oxygen-poor environments where there may be an open wound.18

Bleeding can lead to life-threatening hemorrhage.
Every woman going through a chemical abortion does a considerable amount of bleeding, but when this bleeding is heavy and does not stop, she can be in serious danger if she does not get surgical treatment.19

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3. FDA, 01/2023 Mifepristone Label (see note 1), accessed 5/10/23, Table 4, p. 13 shows risk of failure – defined in terms of “Surgical intervention for ongoing pregnancy” – increased with gestational age. Though that chart showed progression only up to ten weeks, abortion pill advocates led by Elizabeth G. Raymond and Daniel Grossman have advocated a “No-Test” chemical abortion protocol (see note 1) which would rely on prescriber and patient estimates based on last menstrual period or LMP. Authors note that “Regardless of the precise GA [gestational age] limit selected, use of the no-test approach will inevitably result in treatment of some fraction of patients whose true GAs exceed 77 days” [already beyond the FDA limit], p. 363. More recent stories and studies show higher rates of complications with a pharmacy or home delivery system. Researchers from Canada looking at nearly 40,000 abortion patients in Ontario between 2017 and 2020 found nearly 10.3% of the chemical abortion patients visiting the emergency room with some concern or complaint (Ning Liu and Joel G. Ray, “Short-Term Adverse Outcomes After Mifepristone-Misoprostol Versus Procedural Induced Abortion,” Annals of Internal Medicine, January 3, 2023, online edition). Notably, Canada has a pharmacy distribution system similar to the one proposed for the United States. An analysis by a former executive at Marie Stopes International found higher rates of complications once the “Pills by Post” program was put in place in the United Kingdom. Kevin Duffy of Percuity found that 5.9% of chemical abortion patients were treated for complications connected to incomplete abortions or “retained products of conception.” Three percent of women there require surgery to deal with incomplete abortions and 2.3% of these patients were treated in National Trust hospitals for hemorrhage. Rates were higher after the institution of the mifepristone mailing program than before (Percuity, 10/27/21, at https://percuity.files.wordpress.com/2021/10/foi-ma-treatment-failure-211027.pdf, accessed 5/10/23). Another source found a jump in ambulance calls in Britain after the “Pills by Post” system went into operation, increasing by more than 50% in some areas, up at least 25% in others (“Home abortion pills spark major review demand as emergency call outs double in some areas,” Daily Express (London), April 25, 2023, found at express.co.uk/news/politics/1762710/home-abortion-pills-call-outs-review-demand, accessed 5/10/23.

4. Alice Cartwright of Advancing New Standards in Reproductive Health (ANSIRH), a research group from the University of California - San Francisco that, among other things, promotes abortion by telemedicine, writes specifically about “How telemedicine can fill the void left by ‘abortion deserts.’” Mashable, May 27, 2018 at https://mashable.com/article/abortion-deserts-telemedicine, accessed 1/24/22. “Abortion deserts” are those places that are more than 100 miles or more from an abortion clinic. Cartwright neglects to mention that a person far from an abortion clinic may also be miles from the closest emergency room, a critical factor should a patient begin to hemorrhage, show signs of infection, or go into shock. For a map showing what an enormous portion of the United States qualifies as “abortion desert,” see Jake Flynn Mogensen, “This Map Depicts Abortion Access Across America and It’s Really Bleak,” Mother Jones, May 15, 2018, at https://www.motherjones.com/politics/2018/05/this-map-depicts-abortion-access-across-america-and-its-really-bleak/, accessed 1/24/22.

5. Chloe Murtagh, Elizabeth Raymond, Beverly Winkoff, et al., “Exploring the feasibility of obtaining mifepristone and misoprostol from the internet,” Contraception, Vol. 97, no. 4 (April 2018), pp. 287-291. Though authors tried to argue that their attempt to order abortion pill kits over the internet showed the method was “feasible,” they found that none of the kits came with any instructions or warnings, none required any prescription, some came with pin pricks in the packaging that may have degraded the products, raising serious questions about the safety and efficacy of these drugs and the responsibility of those selling them.

6. If a healthcare provider does not directly examine a woman or ask for some sort of identification or documentation, he or she cannot be certain that the person they are talking to is actually pregnant or is even the intended patient. Even before the FDA authorized prescription of abortion pills by telemedicine, there were multiple occasions where someone ordered pills that they intended to use on others, e.g., Kevin Murphy, “Abortion-drug dealer pleads guilty, linked to Grand Rapids man accused of poisoning pregnant woman's drink,” Wisconsin Rapids Tribune, March 5, 2020, https://www.wisconsinrapidstribune.com/story/news/2020/03/05/abortion-pill-dealer-ursula-wing-guilty-case-tied-grand-rapids-man/4966488002/. Shereen Siewert, “New York woman convicted of illegally selling abortion drugs in Wisconsin,” Wausau Pilot & Review, July 12, 2020, https://wausaupilotandreview.com/2020/07/12/new-york-woman-convicted-of-illegally-selling-abortion-drugs-in-wisconsin/, accessed 8/7/20.

7. In Stephen L. Fielding, Emme Edmunds and Eric A. Schaff, “Having an Abortion Using Mifepristone and Home Misoprostol: A Qualitative Analysis of Women's Experiences,” Perspectives on Sexual and Reproductive Health, Vol 34, No. 1 (January/February 2002), pp. 34-40, at p. 37, early pioneer Eric Schaff noted early on in the chemical abortion campaign that “Even though a woman may have experienced cramping and bleeding, she cannot know for certain that her abortion is complete until a provider performs either a sonogram or a hormonal pregnancy test.” The FDA concurs. On page 4 of the FDA Mifepristone Label (see note 1), at ‘2.3 Post-treatment Assessment: Day 7 to 14,’ The FDA cautions that “Lack of bleeding following treatment usually indicates failure; however, prolonged or heavy bleeding is not proof of a complete abortion.”
Mifepristone Safety & Efficacy...

8. FDA Mifepristone Label (see note 1) on page 7 says:

Abdominal pain/cramping is expected in all medical abortion patients and its incidence is not reported in clinical studies. Treatment with MIFEPREX and misoprostol is designed to induce uterine bleeding and cramping to cause termination of an intrauterine pregnancy. Uterine bleeding and cramping are expected consequences of the action of MIFEPREX and misoprostol as used in the treatment procedure. Most women can expect bleeding more heavily than they do during a heavy menstrual period.

9. Ob.Gyn. News (1989), No. 24, p. 1, noted that the average blood loss from mifepristone abortion was reported to be 70ml, nearly four times the blood loss from a standard vacuum curette abortion.

10. Planned Parenthood, “In-Clinic Abortion,” says “In-clinic abortions are also much faster than the abortion pill: most in-clinic abortions only take about 5-10 minutes, while a medication abortion may take up to 24 hours to complete” www.plannedparenthood.org/learn/abortion/in-clinic-abortion-procedures, accessed 1/22/22.

11. The FDA’s “Medication Guide” for Mifepristone (part of the Mifepristone Label, note 1) advises women that “medication abortion” is a two-drug, multi-step process where the prostaglandin misoprostol is taken 24 to 48 hours after the first drug, mifepristone. Though cramps and bleeding may ensue within 2-24 hours after taking the misoprostol, the FDA says on page 19 of the guide that “bleeding or spotting can be expected for an average of 9 to 16 days and may last up to 30 days.”

12. The FDA’s “Medication Guide” for Mifepristone (part of the Mifepristone Label, note 1) says on page 17 that “About 2 to 7 out of 100 women taking Mifeprex will need a surgical procedure because the pregnancy did not completely pass from the uterus or to stop bleeding.” Even proponents trumpeting newer protocols admit high failure rates. UCSF Health’s “Aspiration Versus Medication Abortion,” at ucsfhealth.org/education/aspiration-versus-medication-abortion (accessed 5/10/23) says 3-5% require surgery or an “additional aspiration procedure due to ongoing pregnancy, prolonged or excessive bleeding, or preference.”

13. “Table 4” of the FDA’s 01/2023 Label for Mifepristone (see note 1) notes that efficacy diminishes each week after 49 days LMP.


15. “Blood and Tears,” Newsweek, 9/17/85. A 21-year old told Louise Levanthes, of Health (Jan/Feb 1995) “When I looked at it, it had two dark spots like eyes and a little skeleton not quite formed....I haven't talked about it to anyone. I feel quite empty.” See also the Endowment for Human Development (EHD) website on fetal development at 8 weeks at ehd.org/science_main.php?level=i#th9, accessed 1/26/22. EHD lists a number of features that would be present and visible in the child’s eighth week (10 weeks LMP). At this stage of development, the child would be more than an inch tall from the crown of the head to his or her rump and would have a little skeleton not quite formed....

16. FDA, “Mifepristone U.S. Post-Marketing Adverse Events Summary through 6/30/2022,” Reference ID: 5075481, available at https://www.fda.gov/media/164351/download, accessed 5/10/23. Though the FDA stopped soliciting information on complications and non-lethal events in March 2016, the post-marketing report still records not only 28 deaths which include eight cases of death associated with Clostridium sordellii, another with Clostridium perfringens, two cases of ruptured ectopic pregnancy, cases of hemorrhage, overdose, etc., but also hundreds of non-lethal infections, blood loss requiring transfusions, and nearly a hundred cases of ectopic pregnancy. The FDA also notes more than a dozen deaths from other countries which involved bacterial sepsis, hemorrhage, heat attack, and other cases. Reported U.S. complication rates have been lower than those reported by European governments with nationalized medical systems and centralized health registrations which are thought to obtain more complete, accurate counts. A Finnish study from 2009 (M. Niinimäki, et al., “Immediate complications after medical compared with surgical termination of pregnancy,” Obstetrics & Gynecology, Vol 114, No. 4 (October 2009), pp. 1281-89) showed that 20% of chemical abortion patients experienced adverse events (largely a result of a higher rate of hemorrhage of 15.6%).

17. The FDA’s label for mifepristone (see note 1) notes at “5.4 Ectopic Pregnancy” that “Healthcare providers should remain alert to the possibility that a patient who underwent a medically induced abortion may have an ectopic pregnancy.” The presence of an ectopic pregnancy may have been missed even if the patient underwent ultrasonography prior to being prescribed MIFEPREX” (p. 6).

Atypical Presentation of Infection. Patients with serious bacterial infections and sepsis can present without fever, bacteremia or significant findings on pelvic examination. Very rarely, deaths have been reported in patients who presented without fever, with or without hemoconcentration, and general malaise. A high index of suspicion is needed to rule out serious infection and sepsis.

19. In addition to recorded deaths and serious adverse events associated with bleeding in the FDA’s 6/30/21 Mifepristone U.S. Postmarketing Adverse Events Summary mentioned and linked in note 16 above, there are multiple warnings about the possibility of hemorrhage on the FDA’s 01/2023 label for Mifeprex (mifepristone), including a special “black box” warning stating the following:

WARNING: SERIOUS AND SOMETIMES FATAL INFECTIONS OR BLEEDING...
Serious and sometimes fatal infections and bleeding occur very rarely following spontaneous, surgical, and medical abortions, including following MIFEPRex use...
* Bleeding. Prolonged heavy bleeding may be a sign of incomplete abortion or other complications and prompt medical or surgical intervention may be needed.