Drug Regulatory Agency Warnings on Psychiatric drugs and Violence

There have been 22 drug regulatory agency warnings from five countries and the European Union, on psychiatric drugs causing violence, hostility, aggression, psychosis, mania and homicidal ideation. These are as follows:


2. **United States, March 22, 2004:** The FDA issued a Public Health Advisory on antidepressants stating: “Anxiety, agitation, panic attacks, insomnia, irritability, hostility, impulsivity, akathisia [severe restlessness], hypomania [abnormal excitement, mild mania] and mania [psychosis characterized by exalted feelings, delusions of grandeur and overproduction of ideas], have been reported in adult and pediatric patients being treated with antidepressants.” Source: “WORSENING DEPRESSION AND SUICIDALITY IN PATIENTS BEING TREATED WITH ANTIDEPRESSANT MEDICATIONS,” FDA Public Health Advisory, 22 Mar. 2004.

3. **United States, October 1995:** The U.S. Drug Enforcement Administration (DEA) said Ritalin use could lead to addiction and that “psychotic episodes, violent behavior and bizarre mannerisms had been reported” with its abuse. Source: “Methylphenidate,” U.S. Drug Enforcement Administration (DEA), October 1995.

4. **United States, June 28, 2005:** The FDA announced labeling changes for Concerta and other methylphenidate (Ritalin) products (stimulants) to include, “psychiatric events such as visual hallucinations, suicidal ideation, psychotic behavior, as well as aggression or violent behavior.” Source: “Statement on Concerta and Methylphenidate for the June 30 PAC”, Food and Drug Administration (FDA), June 2005.

5. **Canada, February 2006:** Health Canada approved a new warning label for **Paxil** that stated: “A small number of patients taking drugs of this type may feel worse instead of better. For example, they may experience unusual feelings of agitation, hostility or anxiety, or have impulsive or disturbing thoughts, such as thoughts of self-harm or harm to others.“ Health Canada required Paxil’s product information to detail a list of “rare” side effects, including delusions, hostility, psychosis, and psychotic depression. Source: Kate Jaimet, “‘I’ve learned a lesson in the worst way possible’: What drove a loving father to kill his son?,” Ottawa Citizen, 27 Aug. 2006.

6. **Canada, June 03, 2004:** Health Canada issued an advisory to the public that stated people taking antidepressants at any age are at greater risk of behavioral or emotional changes including self-harm or harm to others. The advisory said, “A small number of patients taking drugs of this type may feel worse instead of better…. For example, they may
experience unusual feelings of **agitation**, **hostility** or **anxiety**, or have **impulsive or disturbing thoughts that could involve self-harm or harm to others.**" Source: Jirina Vlk, “Health Canada advises Canadians of stronger warnings for SSRIs and other newer anti-depressants,” Health Canada, 2004-31, June 3, 2004.


**8. European Union, August 19, 2005:** The Commission of the European Communities issued the strongest warning against child **antidepressant** use as recommended by Europe’s Committee for Medicinal Products for Human Use (CHMP) stating that the drugs were shown to cause suicidal behavior including suicide attempts and suicidal ideation, as well as **aggression and hostility (predominantly aggression, oppositional behavior and anger)** and/or related behavior. Source: Commission of the European Communities Commission Decision concerning the placement on the market, under Article 21 of the Directive 2001/83/EC of the European Parliament and of the Council, Brussels 19-VIII-2005, C (2205) 3256.

**9. Australia, February 2009:** The Australian Therapeutic Goods Administration placed a boxed warning (the strongest warning) onto the ADHD psychostimulant drug methylphenidate (**Concerta and Ritalin**) for drug dependence. It warns that chronic abuse of methylphenidate can lead to a marked tolerance and psychological dependence with varying degrees of **abnormal behavior** and **frank psychotic episodes** can also occur. Source: “Boxed Warning, Contraindications and strengthened Precautions for Methylphenidate,” Janssen-Cilag, February 2009.

**10. Australia, December 2004:** The Australian Therapeutic Goods Administration published an Adverse Drug Reactions Bulletin recommending that any use of SSRI **antidepressants** in children and adolescents should be carefully monitored for the emergence of suicidal ideation and that there was an increase in adverse psychiatric events of suicide, **self-harm**, **aggression** and **violence**. Source: “Use of antidepressants in children and adolescents,” The Australian Therapeutic Goods Administration (TGA) published an Adverse Drug Reactions Bulletin, Vol 23, No. 6, Dec. 2004, p. 22.

**11. United States, July 01, 2009:** The FDA required the manufacturers of the smoking cessation aids varenicline (Chantix) and bupropion (Zyban, aka the antidepressant **Wellbutrin**) to add new Boxed Warnings and develop patient Medication Guides highlighting the risk of serious neuropsychiatric symptoms in patients using these products. These symptoms include changes in behavior, **hostility**, **agitation**, depressed mood, suicidal thoughts and behavior, and attempted suicide. Source: “Information for Healthcare Professionals: Varenicline (marketed as Chantix) and Bupropion (marketed as Zyban, Wellbutrin, and generics),” FDA, July 1, 2009.

13. **Australia, December 2008:** The Australian Adverse Drug Reactions Bulletin advised that the psychostimulant Modafinil has been reported to cause serious adverse skin and psychiatric reactions including anxiety, hallucination, aggression, and mania. Source: Adverse Drug Reactions Advisory Committee, Australian Adverse Drug Reactions Bulletin, Vol. 27, No. 6, December 2008.


15. **United States, September 2007:** The Vice President of Medical Services at the drug company Cephalon sent out a letter to health care professionals informing them of new warnings for the company’s psychostimulant Provigil. The letter stated that there are now “Warnings regarding serious rash, including Stevens Johnson Syndrome [a life-threatening condition affecting the skin] and hypersensitivity reactions, and psychiatric symptoms (including anxiety, mania, hallucinations, and suicidal ideation).” Source: Jeffrey M. Dayno, M.D., “Dear Healthcare Professional,” Cephalon, September 2007.

16. **United States, February 21, 2007:** The FDA directed ADHD drug manufacturers to distribute “patient friendly” guides to consumers warning about serious psychiatric and cardiovascular problems, including stroke, heart attack, sudden death and psychotic reactions caused by ADHD drugs. The psychiatric adverse events included hearing voices, becoming suspicious for no reason, or becoming manic, even in patients who did not have previous psychiatric problems. Source: “FDA Directs ADHD Drug Manufacturers to Notify Patients about Cardiovascular Adverse Events and Psychiatric Adverse Events,” FDA News, February 21, 2007.

17. **United States, August 21, 2006:** The FDA said that ADHD drug manufacturers have to strengthen their warning labels to warn that the drugs can cause suppression of growth, psychosis, bipolar illness, aggression, and ‘serious’ cardiovascular side effects, including misuse possibly leading to sudden death from heart attacks and strokes. Source: “UPDATE 2-US FDA calls for new warnings on ADHD drugs”, Reuters, August 21, 2006.

18. **European Union, April 25, 2005:** The European Medicines Agency’s scientific committee, the Committee for Medicinal Products for Human Use, concluded that Prozac-type antidepressants were associated with increased suicide-related behavior and hostility in young people. It recommended the inclusion of strong warnings across the whole of the
European Union to doctors and parents about these risks and that the drugs should not be used in children and adolescents in off label situations. Source: “EU calls for tougher warnings on antidepressants for kids” News-Medical.net April 25, 2005.

19. **United Kingdom, September 21, 2004**: The British Healthcare Products Regulatory Authority issued guidelines that children should not be given most SSRI antidepressants because of clinical trial data showing an increase rate of harmful outcomes, including **hostility**. Source: “Antidepressant aggression concern,” BBC News, 21 Sept. 2004.

20. **European Union, April 22, 2004**: The European Agency for the Evaluation of Medicinal Products issued a press release stating that, according to clinical trials, **Paroxetine** (**Paxil** in the U.S.) could cause suicidal behavior and **hostility** in children. It recommended that Paroxetine not be used in children and recommended that young adults be observed carefully for signs and symptoms of suicidal behavior or **hostility**. Source: “European Agency for the Evaluation of Medicinal Products: Committee for Proprietary Medicinal Products 20-22 April 2004″ EMEA, The European Agency for the Evaluation of Medicinal Products, Press Release April 2004.


22. **United States, May 2007**: The FDA published a warning on the psychostimulant **Desoxyn** stating that the drug could cause sudden death with pre-existing structural cardiac abnormalities or other serious heart problems, psychiatric adverse events including **aggression** and the **emergence of new psychotic or manic symptoms**, long-term suppression of growth, seizures, visual disturbance, as well as serious cardiovascular adverse event. Source: Food and Drug Administration (FDA), “Detailed View: Safety Labeling Changes Approved By FDA Center for Drug Evaluation and Research (CDER)”. FDA MedWatch, May 2007.