Report for Congress

Received through the CRS Web

Human Cloning

Updated May 2, 2002

Judith A. Johnson Specialist in Life Sciences Domestic Social Policy Division

Human Cloning

Summary

On November 25, 2001, a Massachusetts company, Advanced Cell Technology (ACT), announced that it had created the world's first cloned human embryos. The cloned embryos survived only for a few hours. ACT has indicated that it intends to use such embryos to derive stem cells in producing new therapies for diseases like diabetes and Parkinson's disease. The possibility of using cloning technology not just for therapeutic purposes but also for reproducing human beings raises profound moral and ethical questions.

President Bush announced in August 2001 that for the first time federal funds will be used to support research on human embryonic stem cells, but funding will be limited to "existing stem cell lines." President Bush indicated that he is strongly opposed to human cloning, and that federal funds will not be used for the cloning of human embryos for any purpose, including stem cell research. The Bush Administration established The President's Council on Bioethics on November 28, 2001, "to consider all of the medical and ethical ramifications of biomedical innovation." In January 2002 the 18-member Council held its first meeting and Chairman Leon Kass announced that the first topic to be addressed would be human cloning. At the second (February 2002) and third (April 2002) meetings the discussion of human cloning and stem cell research was continued.

On January 18, 2002, the National Academies released its report entitled *Scientific and Medical Aspects of Human Reproductive Cloning*. The panel recommends that the U.S. ban human reproductive cloning that is aimed at creating a child. It suggests the ban should be legally enforceable and carry substantial penalties rather than rely on voluntary actions. It should be reconsidered within 5 years, but only if compelling new data on safety and efficacy are presented and a national dialogue on the social and ethical issues suggests that a review is warranted. However, the panel concluded that research using cloning procedures to produce stem cells should be permitted because of the considerable potential for developing new therapies and advancing biomedical knowledge. This position is in agreement with a previous National Academies' report entitled *Stem Cells and the Future of Regenerative Medicine* which was released on September 11, 2001.

On July 31, 2001, the House passed H.R. 2505 by a vote of 265-162. The bill would ban the process of human cloning as well as the importation of any product derived from an embryo created via cloning. The provisions mean that cloning could not be used for reproductive purposes or for research on therapeutic purposes, which has implications for stem cell research. The Senate is expected to debate the various legislative proposals concerning human cloning in May 2002. S. 1899 (Brownback) is the companion bill to H.R. 2505. On April 10, 2002, President Bush announced his support for S. 1899 and 40 Nobel Laureates, who are in favor of nuclear transplantation technology for research and therapeutic purposes, announced their opposition to the Brownback bill. Senators Arlen Specter, Dianne Feinstein, Orrin Hatch and Edward Kennedy introduced S. 2439 on April 30, 2002. S. 2439 would prohibit human reproductive cloning while allowing cloning for medical research purposes, including stem cell research. This report will be updated as needed.

Contents

Background			1
Ethical and Social Issues			2
Brief History of Federal Policy Involving Human Embryo Research			3
Bush Administration Policy Regarding Human Embryo Research			5
Legislation			

Human Cloning

Background

The term "cloning" is used by scientists to describe many different processes that involve making copies of biological material, such as a gene, a cell, a plant or an animal. The cloning of genes, for example, has led to new treatments developed by the biotechnology industry for diseases such as diabetes and hemophilia. In the context of this report, a human embryo produced via cloning involves the process called somatic¹ cell nuclear transfer (SCNT). In SCNT, the nucleus of an egg is removed and replaced by the nucleus from a mature body cell, such as a skin cell. In cloning, the embryo is created without sexual reproduction.

Concern over the possibility of producing a human clone increased with the announcement on February 24, 1997, that scientists in Scotland had used SCNT in 1996 to produce the first cloned adult mammal, Dolly, the sheep. Scientists at the Roslin Institute in Edinburgh removed the nucleus from a sheep egg and replaced it with the nucleus of a mammary gland cell from an adult sheep. The resulting embryo was then transferred to the uterus of a surrogate sheep. A total of 277 such embryos were transferred, but only one lamb was born.² Analyses of Dolly's genetic material confirmed that she was derived from the sheep mammary cell. Proponents maintain that cloning could be used for a number of significant agriculture applications, including the improvement of livestock.

On November 25, 2001, Advanced Cell Technology (ACT) of Massachusetts announced that it had created the world's first human embryos produced via cloning; the results were published the following day in an electronic journal.³ ACT used two techniques to produce human embryos — SCNT and a second process called parthenogenesis. ACT researchers obtained eggs from seven women, ages 24 to 32, who were paid \$3000 to \$5000. In the SCNT approach, scientists removed the nucleus from 19 eggs and replaced it with a nucleus from another adult cell. For 11 of the eggs, the nucleus came from a skin cell, for the remaining eight eggs, from cells which cling to the egg and are called cumulus cells. None of the eggs that received a skin cell nucleus divided; seven of the eggs with the cumulus cell nucleus began to divide. Two embryos divided into four cells each, and one embryo divided into six cells before division stopped. In parthenogenesis, an egg cell is treated with

¹ A somatic cell is a body cell, as opposed to a germ cell, which is an egg or sperm cell.

² Wilmut, I., et al. Viable Offspring Derived From Fetal and Adult Mammalian Cells. *Nature*, v. 385, February 27, 1997. p. 810-813.

³ Cibelli, J.B., et al. Somatic Cell Nuclear Transfer in Humans: Pronuclear and Early Embryonic Development. *Journal of Regenerative Medicine*, v. 2, November 26, 2001. p. 25-31.

chemicals causing it to divide without being fertilized by a sperm. ACT exposed 22 human eggs to the chemicals. After 5 days, six eggs had matured into a larger mass of cells before division stopped. None of the embryos developed by ACT through cloning divided sufficiently to produce stem cells.

The stated goal of ACT's work is not to produce a cloned human baby (which requires implantation of the cloned embryo into a woman's uterus), but human embryonic stem cells.⁴ Other research groups have derived stem cells from mice and cattle using SCNT. ACT intends to derive stem cells from human embryos to develop new therapies for diseases such as diabetes and Parkinson's disease. Some scientists believe that stem cells transplanted into a patient could treat disease or injury by replacing damaged tissue. If the cell nucleus used in SCNT is from the patient, the stem cells would be genetically identical to the patient, recognized by the patient's immune system, and avoid any tissue rejection problems that could occur in other stem cell therapeutic approaches. Because of this, many scientists believe the SCNT technique may provide the best hope of eventually treating patients using stem cells for tissue transplantation. A California biotechnology company, Geron Corporation, is also working on stem cells created via SCNT.⁵

Ethical and Social Issues

The possibility of using cloning technology not just for therapeutic purposes but also for reproducing human beings raises profound moral and ethical questions. In response to the creation of Dolly and concerns about the potential application of cloning humans, on February 24, 1997, President Clinton asked the National Bioethics Advisory Commission⁶ (NBAC) to review the ethical and legal issues associated with the use of cloning technology; NBAC reported its findings and recommendations on June 9, 1997.⁷ NBAC recommended a continuation of the moratorium on the use of federal funding in the support of any attempt to create a child by SCNT, and an immediate request to all non-federally funded investigators to comply voluntarily with the intent of the federal moratorium. NBAC also recommended that federal legislation be enacted, with a 3- to 5- year sunset clause, to prohibit anyone from attempting to create a child through the use of SCNT in a research or clinical setting. The NBAC found it morally unacceptable to attempt to clone humans for the purpose of human reproduction because scientific data from

[http://www.whitehouse.gov/news/releases/2001/08/20010809-2.html].

⁴ For more information about stem cells, see CRS Report RL31015, *Stem Cell Research*, by Judith A. Johnson.

⁵ Weiss, R. Embryo Work Raises Spector of Human Harvesting. *Washington Post*, June 14, 1999. p. A01.

⁶ NBAC was established by Presidential Executive Order 12975 on October 3, 1995, to provide guidance to federal agencies on the ethical conduct of current and future human biological and behavioral research. A September 16, 1999 executive order extended the NBAC charter until October 2001. NBAC has been replaced by the President's Council on Bioethics, which was described by the Bush Administration in its August 9, 2001 policy decision on human embryonic stem cell research. The President's remarks on embryonic stem cell research are available at:

⁷ National Bioethics Advisory Commission. *Cloning Human Beings*. June 1997.

animal experiments indicate the method is not safe for mother or baby. In addition to concerns about physical safety, the NBAC report pointed out that SCNT raises issues about the individuality, autonomy, objectification and kinship of the resulting children.

Cloning, if allowed for human reproduction, could affect society's perception of what it means to be a human being. Uncertainties over a cloned individual's personal uniqueness or freedom to create one's own identity may haunt him or her. Relatives or friends could have specific expectations regarding the cloned individual's talents and abilities. Others might ill treat or discriminate against a cloned individual. Some worry that cloning would lead to diminished respect for human life in general, and for cloned individuals in particular, since the cloned person might simply be replaced with another clone. Others point out, however, that this altered perception does not occur today with identical twins, who are naturally produced clones. Cloning human embryos also raises difficult questions about the rights of parents to control their own embryos and other issues concerning reproductive rights and privacy. Some observers believe that it would be ethical to clone human embryos to help infertile couples conceive. Lastly, the possibility of human cloning is offensive to the religious and other deeply held beliefs of many people.

On January 7, 1998, a Chicago scientist, Dr. Richard Seed, announced his intention to clone a human being. In response, bills were introduced in Congress that would have banned human cloning indefinitely or imposed a moratorium. The legislation was opposed by a number of medical organizations, the biotechnology industry and many scientists and was not enacted. Others expressing an interest in reproductive cloning include: (1) Clonaid, a company directed by chemist Brigitte Boisselier and formed by the Raelians, a group that believes humans are descendants of extraterrestrials and that cloning can allow humans to become immortal; and, (2) Panos Zavos, of the University of Kentucky, and Severino Antinori, director of a fertility clinic in Rome, who are working together to help infertile couples have children via cloning. In April 2002, there were unconfirmed reports in the media that both Clonaid and Severino Antinori had implanted cloned human embryos in women. Dr. Severino claims there are 3 such pregnancies of 6 to 9 week duration; 2 in Russia and 1 in an Islamic state. His claim has been disputed by his former partner, Panos Zavos. The Clonaid group would not reveal any further details about their cloning attempts.

Brief History of Federal Policy Involving Human Embryo Research

Currently no U.S. laws or regulations would prohibit all cloning research. However, federal funding of *any* type of research involving human embryos, starting with *in vitro* fertilization (IVF) then later cloning and stem cells, has been blocked by various policy decisions dating back 25 years. Following the birth of the first IVF baby, Louise Brown, in July 1978, the Ethics Advisory Board (EAB) was tasked with considering the scientific, ethical, legal, and social issues surrounding human IVF.⁸ The EAB released its report on May 4, 1979, which found that IVF research was acceptable from an ethical standpoint and could be supported with federal funds. The EAB's recommendations were never adopted by HHS, the EAB was dissolved in 1980, and no other EAB was ever chartered. Because federal regulations that govern human subject research (45 CFR 46) stipulated that, at the time, federally supported research involving human IVF must be reviewed by an EAB, a so-called "de facto moratorium" on human IVF research resulted. Other types of embryo research ensuing from the development and use of IVF, such as cloning and stem cells, were therefore also blocked. The de facto moratorium was lifted with the enactment of the National Institutes of Health (NIH) Revitalization Act of 1993 (P.L. 103-43, Section121(c)) which nullified the regulatory provision (45 CFR 46.204(d)) requiring EAB review of IVF proposals.

In response, the NIH established the Human Embryo Research Panel to assess the moral and ethical issues raised by this research and develop recommendations for NIH review and conduct of human embryo research. The NIH Panel released a report providing guidelines and recommendations on human embryo research in September 1994. It recommended that some areas of human embryo research be considered for federal funding, including SCNT, stem cells and (under certain limited conditions) embryos created solely for the purpose of research.⁹ The NIH Panel also identified areas of human embryo research it considered to be unacceptable, or to warrant additional review. It determined that certain types of cloning¹⁰ without transfer to the uterus warranted additional review before the Panel could recommend whether the research should be federally funded. However, the Panel concluded that federal funding for such cloning techniques followed by transfer to the uterus should be unacceptable into the foreseeable future. The Panel's report was unanimously accepted by the NIH Advisory Committee to the Director (ACD) on December 2, 1994.

After the ACD meeting on December 2, 1994, President Clinton directed NIH not to allocate resources to "support the creation of human embryos for research purposes." The President's directive did not apply to research involving so-called "spare" embryos, those that sometimes remain from clinical IVF procedures performed to assist infertile couples to become parents. Nor did it apply to human parthenotes, eggs that begin development through artificial activation, not through fertilization. Following the Clinton December 2, 1994 directive to NIH, the agency

⁸ The EAB was created in 1978 by the Department of Health Education and Welfare (HEW), the forerunner of the Department of Health and Human Services (HHS). The EAB was formed at the recommendation of the National Commission for the Protection of Human Subjects of Biomedical and Behavioral Research. The National Commission operated from 1974 to 1978 and issued 10 reports, many of which formed the basis of federal regulations for research involving human subjects (45 CFR 46).

⁹ National Institutes of Health. *Report of the Human Embryo Research Panel*, September 27, 1994.

¹⁰ These were **blastomere separation**, where a two- to eight-cell embryo is treated causing the cells (blastomeres) to separate; and, **blastocyst division**, in which an embryo at the more advanced blastocyst stage is split into two.

proceeded with plans to develop guidelines to support research using spare embryos. However, these plans were halted on January 26, 1996, with the enactment of P.L. 104-99 which contained a rider that affected FY1996 funding for NIH. The rider prohibited HHS from using appropriated funds for the creation of human embryos for research purposes or for research in which human embryos are destroyed. This same rider, often referred to as the Dickey amendment, has been attached to the Labor, HHS and Education Appropriations Acts for FY1997 through FY2002.¹¹ Current language, Section 510 of the FY2002 Labor, HHS and Education Appropriations Act, prohibits HHS from using FY2002 appropriated funds for:

(1) the creation of a human embryo or embryos for research purposes; or,

(2) research in which a human embryo or embryos are destroyed, discarded, or knowingly subjected to risk of injury or death greater than that allowed for research on fetuses in utero under 45 CFR 46.208(a)(2) and Section 498(b) of the Public Health Service Act (42 U.S.C. 289g(b)). For purposes of this section, the term "human embryo or embryos" includes any organism, not protected as a human subject under 45 CFR 46 ... that is derived by fertilization, parthenogenesis, cloning, or any other means from one or more human gametes [sperm or egg] or human diploid cells.

One month after the Dolly announcement, on March 4, 1997, President Clinton sent a memorandum to the heads of all executive departments and agencies making it "absolutely clear that no federal funds will be used for human cloning." This action extended the congressional ban beyond HHS to all federally supported research. Clinton also urged the private sector to adopt a voluntary ban on the cloning of human beings. The *NIH Guidelines on Stem Cell Research*, published by the Clinton Administration in August 2000, would not have funded research in which: human stem cells are used for reproductive cloning of a human; human stem cells are *derived* using SCNT; or, human stem cells that were derived using SCNT are *utilized* in a research project.

Bush Administration Policy Regarding Human Embryo Research

On August 9, 2001 President Bush announced that for the first time federal funds will be used to support research on human embryonic stem cells, but funding will be limited to "existing stem cell lines." In the speech, President Bush stated that he strongly opposes human cloning. Although not mentioned specifically in the August 9 speech, a fact sheet on the White House website states that federal funds

¹¹ The original rider, introduced by Representative Jay Dickey, is in Section 128 of P.L. 104-99; it affected NIH funding for FY1996 contained in P.L. 104-91. For subsequent fiscal years, the rider is found in Title V, General Provisions, of the Labor, HHS and Education Appropriations Acts in the following public laws: FY1997, P.L. 104-208; FY1998, P.L. 105-78; FY1999, P.L. 105-277; FY2000, P.L. 106-113; FY2001, P.L. 106-554; and FY2002, P.L. 107-116.

will not be used for "the cloning of human embryos for any purpose."¹² In his speech, President Bush announced his intention to name a President's council, chaired by Dr. Leon Kass of the University of Chicago, "to consider all of the medical and ethical ramifications of biomedical innovation." The President's Council on Bioethics, was established for a period of up to 2 years by Executive Order 13237 on November 28, 2001. The White House announced the other 17 members of the council on January 16, 2002.

The first meeting of the President's Council on Bioethics was held on January 17-18, 2002, in Washington, D.C.¹³ Dr. Kass announced that the first topic to be addressed by the Council would be human cloning. At the Council's second meeting, the terminology of cloning was discussed in order to reach a consensus on the terms used to describe the two types of cloning: reproductive vs. therapeutic or research cloning. Although all Council members voted in opposition to reproductive cloning, they could not come to an agreement on articulating the precise nature of their objection, whether solely on safety grounds or which of the various moral objections were most important. On the issue of therapeutic cloning, what the Council prefers to call research cloning, the Council also could not come to agreement. Dr. Kass proposed that the Council's final report should reflect both the arguments supporting cloning for the purpose of medical treatment and those against. He asserted that the report should also provide the soundest arguments for each position and indicate how many Council members supported each viewpoint.

The third meeting of the Council was held on April 25 and 26, 2002. The Council heard presentations on the scientific and therapeutic promise of embryonic stem cells from John Gearhart of Johns Hopkins University and the potential of adult stem cells from Catherine Verfaillie of the University of Minnesota. In an informal vote, almost half of the 18 members of the Council voiced their support for the therapeutic use of human cloning. A final report by the Council on human cloning is expected by the end of the summer of 2002.¹⁴

A year ago in March 2001, the Food and Drug Administration (FDA) sent letters to the research community stating that the creation of a human being using cloning is subject to FDA regulation under the Public Health Service Act and the Food, Drug and Cosmetic Act.¹⁵ FDA stated that such research could only occur when an investigational new drug application (IND) is in effect. Some legal scholars believe that there is no legal basis for the regulation of cloning by FDA.¹⁶ They find little

¹² The White House Fact Sheet on embryonic stem cell research is available at: [http://www.whitehouse.gov/news/releases/2001/08/20010809-1.html].

¹³ A transcript of the first meeting and papers developed by staff for discussion during the meeting can be found at [http://www.bioethics.gov].

¹⁴ Shehzad, N. President's Bioethics Advisors Resolve to Avoid "Media Circus" as They Work on Cloning Advice." *Washington Fax*, April 30, 2002.

¹⁵ The FDA position statement and letters to the research community are available at [http://www.fda.gov/cber/genetherapy/clone.htm].

¹⁶ Weiss, R. Legal Barriers to Human Cloning May Not Hold Up. *Washington Post*, May (continued...)

evidence to support FDA's position that cloned human embryos are "drugs." However, the biotechnology industry and the American Society for Reproductive Medicine believe FDA has the authority to regulate cloning and legislation is unnecessary because FDA regulation is preferred to any new action by Congress.¹⁷

On January 18, 2002, the National Academies released its report entitled *Scientific and Medical Aspects of Human Reproductive Cloning*.¹⁸ The panel recommends that the U.S. ban human reproductive cloning that is aimed at creating a child. Based on the results of animal cloning experiments, the panel is concerned for the safety of both the woman and the fetus and judged the procedure to be too dangerous for use in humans at the present time. It recommends that the ban should be legally enforceable and carry substantial penalties rather than be based on voluntary actions. It should be reconsidered within 5 years, but only if compelling new data on safety and efficacy are presented and a national dialogue on the social and ethical issues suggests that a review is warranted. However, the panel concluded that research using SCNT to produce stem cells should be permitted because of the considerable potential for developing new therapies and advancing biomedical knowledge. This position is in agreement with a previous National Academies' report entitled *Stem Cells and the Future of Regenerative Medicine* which was released on September 11, 2001.¹⁹

Legislation

On July 19, 2001, the House Judiciary Subcommittee on Crime approved the Human Cloning Prohibition Act of 2001, H.R. 2505 (Weldon), by voice vote. The bill would ban the process of human cloning as well as the importation of any product derived from an embryo created via cloning. The provisions mean that cloning could not be used for reproductive purposes or for research on therapeutic purposes, which has implications for stem cell research. The bill includes a criminal penalty of imprisonment of not more than 10 years and a civil penalty of not less than \$1 million.

On July 24, 2001, the House Judiciary Committee approved H.R. 2505 by a vote of 18 to 11 and defeated a substitute measure by a vote of 11 to 19. The substitute was identical to H.R. 2608 (Greenwood), which would ban *only* human reproductive cloning; the ban would sunset after 10 years. H.R. 2608 has the same criminal and civil penalties as H.R. 2505 when cloning is used "with the intent to initiate a

[http://www.nap.edu/catalog/10285.html?onpi_topnews_011802].

¹⁶ (...continued)

^{23, 2001.} p. A1.

¹⁷ Ibid.

¹⁸ The National Academies are the National Academy of Sciences, the National Academy of Engineering, the Institute of Medicine, and the National Research Council. The report on human cloning is available at:

¹⁹ The National Academies' report on stem cell research is available at: [http://www.nap.edu/catalog/10195.html?onpi_topnews_091101].

pregnancy." The Bush Administration announced its support for H.R. 2505 on July 24, 2001.

On July 31, 2001, the House passed H.R. 2505 by a vote of 265-162. Prior to the vote on H.R. 2505, the House defeated a substitute amendment, H.Amdt. 285, which is identical to H.R. 2608, by a vote of 178 to 249. During debate, supporters of H.R. 2505 argued that a partial ban on human cloning, such as H.R. 2608, would be impossible to enforce. Critics of H.R. 2505 argued that SCNT creates a "clump of cells" rather than an embryo, and that the measure would curtail medical research and prevent Americans from receiving life-saving treatments created overseas.

On December 3, 2001, the Senate considered an amendment proposed by Senator Lott that would have imposed a 6-month moratorium on all human cloning research; an attempt to attach the amendment to a bill (H.R. 10) on pension contribution limits failed (Senate Roll Call vote 344).

The Senate is expected to debate the various legislative proposals concerning human cloning before the Memorial Day recess in late May.²⁰ S. 1899 (Brownback), the Human Cloning Prohibition Act of 2001, was introduced on January 28, 2002; it is the companion bill to H.R. 2505. S. 1899 currently has 30 cosponsors and is very similar to S. 790, a bill introduced by Senator Brownback in April 2001. At a White House press briefing on April 10, 2002, President Bush again stated his support for a prohibition on all forms of human cloning and endorsed Senator Brownback's bill.

On the same day as the White House briefing, the American Society for Cell Biology released a statement, signed by 40 Nobel Laureates, in favor of nuclear transplantation technology for research and therapeutic purposes and in opposition to the Brownback bill.²¹ The statement asserts that S. 1899 "would impede progress against some of the most debilitating diseases known to man."

Former President Gerald Ford stated his strong opposition to both H.R. 2505 and S. 1899 in a April 25, 2002, letter to President Bush.²² In the letter, Ford indicates that during his administration, the controversy over recombinant DNA research was "successfully addressed with 'careful thought' and the implementation of safety regulations."²³ Former President Ford "expresses full support for therapeutic cloning, arguing a prohibition of this technology 'would adversely impact scientific research and should not become law."²⁴

²⁰ Glendenning, D. Cloning Bill Options Set For Senate Floor Debate Before Memorial Day, Daschle Aide Says. *Washington Fax*, April 1, 2002.

²¹ The American Society for Cell Biology statement by the 40 Nobel Laureates is available at: [http://www.ascb.org/publicpolicy/Nobelletter.html].

²² Hafner, L. Revised Feinstein/Kennedy Cloning Bill Has Criminal and Civil Penalties, Requires Research Review. *Washington Fax*, May 2, 2002.

²³ Ibid.

²⁴ Ibid.

Senators Arlen Specter, Dianne Feinstein, Orrin Hatch and Edward Kennedy introduced S. 2439 (Specter), the Human Cloning Prohibition Act of 2002, on April 30, 2002. S. 2439 would prohibit human reproductive cloning while allowing cloning for medical research purposes, including stem cell research. According to a press release from Sen. Specter's office, S. 2439 applies Federal ethical regulations on human subject research to nuclear transplantation research, such as review by an ethics board, inclusion of protections for research participants, privacy and informed consent. It would also impose a \$250,000 fine for a violation of these conditions. S. 2439 would impose penalties for violations of the reproductive cloning ban of up to 10 years in prison and a minimum fine of \$1 million. The bill defines human cloning as "implanting or attempting to implant the product of nuclear transplantation into a uterus or functional equivalent of a uterus."²⁵

Some legal scholars believe a ban on human cloning may be unconstitutional because it would infringe upon the right to make reproductive decisions which is "protected under the constitutional right to privacy and the constitutional right to liberty."²⁶ Other scholars do not believe that noncoital, asexual reproduction, such as cloning, would be considered a fundamental right by the Supreme Court. A ban on human cloning research may raise other constitutional issues: scientists' right to personal liberty and free speech. In the opinion of some legal scholars, any government limits on the use of cloning in scientific inquiry or human reproduction would have to be "narrowly tailored to further a compelling state interest."²⁷

²⁵ The Specter press release can be found at: [http://www.senate.gov/~specter/020426.html].

²⁶ Andrews, L. B. Is There a Right to Clone? Constitutional Challenges to Bans on Human Cloning. *Harvard Journal of Law and Technology*, summer 1998. p. 643-680.

²⁷ Ibid., p. 667.