April 1, 2014

State of Connecticut
S.B. 413

sponsored by the Joint Committee on Public Health

Prepared Testimony of Catherine Glenn Foster, Esq.

My name is Catherine Glenn Foster. I serve as litigation counsel in the Washington, DC, office of Alliance Defending Freedom, a non-profit legal organization based in Scottsdale, Arizona, and with offices around the country and world. I have been involved in numerous cases relating to denial of care and have had the opportunity to advise clients and my fellow attorneys about proposed legislation and compliance with state laws. I write today based on my experience as a patient advocate to express my concerns about the constitutionality and wisdom of this bill.

S.B. 413 on Medical Orders for Life-Sustaining Treatment (“MOLST”) would start a trial program under the Department of Public Health to create a new form of medical orders in Connecticut. The new orders would permit individuals to predetermine what forms of treatment they may wish to choose or refuse in a given situation, prior to being in the context of that situation. It sets preferences in stone. While S.B. 413 was proposed with the best of intentions and crafted so as to avoid some of the more obvious pitfalls of similar forms in other states, in the real world this bill would nonetheless carry very dangerous unintended consequences.

Advance directives assume that an individual’s preferences will remain substantially the same throughout their lives, or that the patient will amend their preferences in writing. Advance directives purport to offer patient autonomy, but block the patient from the fully informed consent that would yield true patient autonomy. Life, and medical treatment in particular, are simply more nuanced than can be captured in an advance directive.

Two of the most common complications that can arise after a person commits to an advance directive are misdiagnosis and the person changing their mind. More than 40% of patients with disorders of consciousness are misdiagnosed\(^1\); this rate has not

\(^1\) See, e.g., Martin M. Monti et al., *Willful Modulation of Brain Activity in Disorders of Consciousness*, 362 NEW ENGLAND J. OF MED. 579 (2010) (noting that the rate of misdiagnosis of disorders of consciousness is approximately 40%); K. Andrews et al., *Misdiagnosis of the
changed despite medical advances over the last 15 years.\textsuperscript{2} Overall, it is estimated that up to 15\% of diagnoses are incorrect in most areas of medicine.\textsuperscript{3} In my legal practice I have seen patients who were clearly misdiagnosed, and even communicating, but whose hands were bound by their advanced directive. The results are often tragic.

And even with accurate diagnoses, patients frequently change their minds. The circumstances in which patients are handed an advance directive form vary, but individuals can frequently be coerced into signing the advance directive or pressured into certain choices. This bill tries to reduce that risk by having trained health care professionals present both benefits and risks, but a limited training session cannot adequately protect against real-world abuses. And S.B. 413 allows either the patient or a surrogate – not limited to a medical power of attorney (“MPOA”) – to sign the advance directive.

In the light of day, or over a period of time, nearly half of patients change their mind or beliefs.\textsuperscript{4} This is even more relevant once the person is in the context of a situation where the advance directive may be enforced, as no advance directive could possibly provide for every clinical scenario. When people are sick or near the end of life, they may suffer “depression, hopelessness and fear of loss of autonomy and control.”\textsuperscript{5} But with counseling and caring, and proper treatment,\textsuperscript{6} there is hope for comfort, and all of these factors may affect a person’s wishes.

Connecticut citizens have expressed their concern about many of these issues: misdiagnosis, sloppy procedures on the part of doctors, a reduction in end-of-life options, and pressure from facilities and families on elder or infirm individuals.\textsuperscript{7} They worry that the focus will be on saving money, not saving lives, and their fears are supported by the medical literature.\textsuperscript{8} And hospitals can easily develop a reliance on advance directives to

\textsuperscript{2} See Caroline Schnakers et al., \textit{Diagnostic Accuracy of the Vegetative and Minimally Conscious State: Clinical Consensus Versus Standardized Neurobehavioral Assessment}, 9 BMC NEUROLOGY 35 (2009).


\textsuperscript{4} See, e.g., Terri R. Fried et al., \textit{Changes in Preferences for Life-Sustaining Treatment Among Older Persons with Advanced Illness}, 22 J. GEN. INTERN. MED. 495 (2007) (studying 189 patients over a two-year period and finding that nearly half were inconsistent in their wishes, even among those with stable health, and even among those with an advance directive); see also http://www.kevinmd.com/blog/2012/09/patients-deviate-advance-directives.html.

\textsuperscript{5} http://opinionator.blogs.nytimes.com/2012/10/27/four-myths-about-doctor-assisted-suicide/.


\textsuperscript{8} See, e.g., C.V. Chambers et al., \textit{Relationship of Advance Directives to Hospital Charges in a Medicare Population}, 14 ARCH. INTERN. MED. 541 (1994) (finding that by limiting care to Medicare patients, “an enormous cost savings to society may be realized”).
shield themselves from wrongful-death and other lawsuits when they choose to deny care to a patient in need.

Concerns such as the high rate of inconsistency in patient desire led Rebecca L. Sudore of University of California, San Francisco, and Terri R. Fried of Yale University School of Medicine to declare,

[Adv]ance directives . . . frequently fail to affect the quality of care received at the end-of-life or improve clinicians’ and surrogates’ knowledge of patients’ preferences. [Despite s]ubstantial improvements . . ., many of these efforts continue to be aimed at . . . the traditional objective of making advance decisions – an objective which is fundamentally flawed. 9

They propose instead patient preparation for participation in in-the-moment decisions. 10

In place of S.B. 413, I submit the following recommendations. First, for patients who have elected to complete “Do Not Resuscitate” (“DNR”) and similar orders, issue guidance to healthcare professionals and others recommending they prepare and point to such orders in a way that will be eye-catching (e.g., by placing a flag on patient files or printing the orders on brightly colored paper) and transferable between facilities and providers.

Before any other advance directive bills are considered, there must be studies to appropriately inform the public of issues such as the circumstances under which patients agree to an advance directive, the average time difference between advance directive preparation and implementation, patients who change their mind and why, the role of advance directives in cases of misdiagnosis, health care professionals who do not adhere to advance directives, and other misuse of advance directives. Only then can the legislature and the public participate in an informed debate on the matter.

Finally, and most importantly, health care representatives must be fully empowered as a MPOA. In my experience, the one form of advance directive least subject to abuse and premature ossification is the MPOA. The goal of all these methods of advance directive is patient autonomy, and MPOAs allow the patient to choose a trusted individual and fully express his beliefs and wishes over time through nuanced discussions. When needed, MPOA then allows the agent the needed freedom to use his substituted judgment to advance the patient’s wishes. This view is supported by physicians like Rebecca L. Sudore and Terri R. Fried, who believe that when a patient is not able to participate in a decision, the best way to achieve patient autonomy is through a surrogate.11 Yet in 2006, S.B. 317 (effective October 1, 2006) combined the power of

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10 Id.
11 Id.
the health care agent and the attorney-in-fact for health care decisions, and expanded the
scope and weight of the living will form of advance directive. The most effective way to
ensure patient autonomy and self-determination would be to create a simple, fast way for
individuals to designate a trusted medical power of attorney, and truly enable that agent
to respond to the nuances of belief and of medical treatment that are absent from “check a
box” advance directives.

S.B. 413 is premature. The State of Connecticut should not waste its valuable
resources implementing a pilot program without first fully exploring the potential
ramifications, and should not put patient lives at risk.

As Muriel Gillick, a geriatrician at Harvard Medical School and researcher in
dead-of-life care, wrote in the New England Journal of Medicine, “Despite the prodigious
effort devoted to designing, legislating, and studying of advance directives, the consensus
of medical ethicists, researchers in health care services, and palliative care physicians is
that the directives have been a resounding failure.”12 S.B. 413 should be shelved until we
have the research and analysis that will allow us to fully engage in that discussion.

12 See JEROME GROOPMAN & PAMELA HARTZBAND, YOUR MEDICAL MIND: HOW TO DECIDE
WHAT IS RIGHT FOR YOU (2011).