The Latest from Sparrow Care Network

Education Update
If you were unable to attend the November 3, 2015 SCN educational program “SCN Meets the Market”, you now have the ability to view the recording. You also are able to earn engagement points in the SCN Incentive Program should you choose to participate in a brief quiz after viewing the video. To learn more, please contact Nicole Allen at 517.364.8158 or nicole.allen@sparrow.org

Save the Date!
“Billing & Coding in a Risk-Adjustment Environment”

Tuesday, March 22 6:30 p.m. – 8 p.m.
(Dinner available at 5:30 p.m.)
The James B. Henry Center for Executive Development
Large Amp Room (A-170)
3535 Forest Road, Lansing, MI 48910

Audience
Sparrow Care Network Physicians

Program Agenda
5:30 p.m.            Dinner
6:30 p.m.            Presentation
7:45 p.m.            Q&A
8:00 p.m.            Adjournment

Why Attend:
Accurate coding is essential as health care continues to move from a fee-for-service to a fee-for-value payment model. The changing reimbursement model requires Physicians and staff to document and capture the information needed to improve risk scores as well as reimbursement rates.

This program is a Physician-focused follow-up to our September billing and coding sessions. Local Physicians who attended the September program found it extremely beneficial and have requested the program return for an additional session.

Continuing Medical Education credit is available.

Ms. Laurie Latvis, RN, Director of Provider Outreach BCBSM, will present the following:
- Risk, coding and documentation
- HEDIS Measures and Medicare Star Ratings
- Improving patient satisfaction
- Incentive Programs
And more

To RSVP:
Contact Nicole Allen with your name, email address and clinic information at 517.364.8158 or nicole.allen@sparrow.org

SCN Physician Performance Dashboard Implementation Update

Last fall, SPH Analytics began to grant SCN Physicians access to their Physician Performance Dashboards. At this time, approximately half of all SCN Physicians now have access to their Dashboard.

During this initial implementation phase it is very important for Physicians and Office Managers to login and validate the accuracy of the displayed data.

The Dashboards are an important tool in ensuring SCN’s compliance with the Department of Justice’s requirements for a Clinically Integrated Organization. The Dashboards also provide performance data to Physicians to assist with monitoring and measuring their performance on the SCN Quality Program Initiatives.

Since these Dashboards play such an important role for SCN it is imperative that this data is correct both for SCN’s goal of improving quality and efficiency, as well for your benefit, as this data will be used for Physician incentives.

SCN Leadership is continuing to contact Physicians and Practice Leaders to further the implementation process. If you have any questions, need assistance with logging into your Dashboard, or are unsure of where you may be in the implementation process, please contact Nicole at 517.364.8158 or nicole.allen@sparrow.org

Announcements

Congratulations to Cornerstone Family Practice and SMG St. Johns who have both been chosen as a Michigan Primary Care Transformation Project (MiPCT) top performer in HEDIS measures. These practices have been asked to participate in a “best practice” site visit with MiPCT leaders.

The purpose of the MiPCT practice visit is to:

1) Observe and document best practice to build MiPCT community learning resources
2) Share best practice with MiPCT POs to achieve statewide project metric
3) Develop a repository of best practices to support sharing with key national and state stakeholders including MiPCT payers

Article of Interests

While modern medicine has afforded people longer lives, end of life care has not advanced as quickly. In fact, studies are beginning to show that end of life medical care may be getting worse.

Palliative Care is a field that is often under-used, and misunderstood. Palliative Care experts can assist seriously ill patients with setting and achieving goals in all areas of their lives, and this can better help those patients with pain and anxiety control, ease family burden and even potentially increase the patient’s chance of survival.

Please read the attached article by Atul Gawande regarding end of life care, “Quantity and Quality of Life, Duties of Care in Life-Limiting Illness.”
How To Learn More

If you’d like to learn more about Sparrow Care Network either via conference call or an informational in-office visit, please don’t hesitate to contact SCN leadership.

Questions about Sparrow Care Network?

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For more information on Clinical Integration, please visit: SparrowCareNetwork.org

SparrowTV.org
Quantity and Quality of Life
Duties of Care in Life-Limiting Illness

Atul Gawande, MD, MPH

Everyone dies. Dying today typically involves a period of protracted illness, disability, and intense involvement of medical professionals. Although the experience is woefully understudied, a significant body of evidence is emerging to guide clinicians, health systems, and society toward better practices for people facing serious, life-threatening conditions.

That evidence has shown, importantly, that the amount of suffering that people endure in their last year of life is considerable. Singer et al1 recently reported on the experience of 7,204 adults older than 50 years who died while being followed up as part of a longitudinal study of US health and retirement. The researchers found that, during their last year, 51% of study participants were often troubled by moderate to severe depression, periodic confusion, dyspnea, and incontinence during that time.1 Furthermore, among those who died between 1998 and 2010, none of these symptoms decreased in occurrence during their last year of life, but rather occurrence of pain, depression, and periodic confusion actually increased. Medical care for the symptoms people experience at the end of life does not seem to have gotten better; it may have gotten worse.

It could be argued that the findings simply reflect people following their wishes to trade the quality of their lives for therapies that extend their lives. But this is not the case. In 2014, the Institute of Medicine (IOM) published Dying in America, which included an extensive review of the medical literature on the end of life, including the efficacy of expert palliative care.2 Palliative care is a field dedicated to assisting seriously ill people with setting and achieving goals aside from just survival, which may include control of symptoms, attending to life projects, connecting with loved ones, or other vital objectives. The literature has established that when care is provided with a narrow focus on disease control, without palliative care expertise directed at eliciting these broader goals and tailoring care to include them, patients experience more pain, more anxiety, and more family exhaustion; they receive more nonbeneficial care and more hospitalization; and they do not live longer. Indeed, studies suggest that earlier involvement of palliative care specialists, either through consultation or enrollment in hospice, can produce increased survival.3,4

Another cause of physical suffering and family distress identified in the IOM report is that the majority of people reach the end of life cognitively impaired by illness, treatment, or frailty, and they are unable to make their own decisions about their care. They must rely on a proxy to make their medical choices. Only with advance planning conversations can they ensure alignment between the treatment they receive and their goals and values. But most people do not have these conversations with their clinicians or family members. Treatment therefore often violates their preferences, usually in the direction of undervaluing measures to support the quality of their remaining life.

The picture of care at the end of life that emerges is therefore disturbing. A widespread perception among both the medical profession and the public at large has been that seeking palliative care consultation or hospice services, or even just having advance planning discussions, amounts to “giving up” and is only relevant when people no longer have options for disease-based therapy. This view is incorrect and harmful.

It is also sometimes regarded as uniquely American. However, in this issue of JAMA, dedicated to the topic of death, dying, and the end of life, Bekelman et al5 report findings that suggest otherwise. The authors analyzed data from patients older than 65 years who died with cancer in 7 countries: Belgium, Canada, England, Germany, the Netherlands, Norway, and the United States.5 Among these countries, the United States actually had the lowest proportion of patients whose site of death was the hospital (22% vs up to 51% elsewhere). To be sure, the United States registered the highest rate of intensive care unit admission and chemotherapy administration in the last 180 days of life, indicating a mixed and still complex picture of care at the end of life. High use of intensive care and other technologies is a general characteristic of US health care,6 and the study found that use of chemotherapy in the last 30 days of life declined in the United States, just as in the other countries, to approximately 10%. All of these countries appeared to be making a radical transition away from nearly universal reliance on hospitalization at death for cancer patients, with the United States, perhaps surprisingly, having moved the farthest and fastest away from institutionalization at death. Other research shows the United States now has perhaps the highest level of hospice capacity and use, and the highest likelihood of death at home, in the developed world.7

Data from another study in this issue of JAMA, by Wright et al,8 indicate that this shift has likely been beneficial. The authors report survey data from family members of 1,146 elderly patients who died with advanced lung or colorectal cancer. Patients who died in the hospital proved the least likely to be reported to have excellent quality of care near death (just 41% did) or to die where they had wished to (only 18% preferred to die in the hospital and did so). By contrast, patients who had been enrolled in hospice for at least 3 days had the best reported quality of life near death and best alignment of care with their wishes.
However, only half of the patients in this multiregional study received 3 or more days of hospice. The authors’ conclusion was therefore, reasonably, that efforts should promote earlier hospice enrollment and avoid hospital death. However, this interpretation is incomplete. The results also suggest that palliative care for hospitalized patients must improve. There is no intrinsic reason that the quality of care for the dying should be poorer in hospitals than elsewhere. Indeed, as Angus and Truog point out in their Viewpoint in this issue of JAMA, even intensive care units can provide critical resources for improvement in the quality of life of dying patients.6

Furthermore, care in out-of-hospital settings can still be inadequate. As Wright et al noted, more than 40% of bereaved family members reported less-than-excellent quality of care near death for patients who received more than 3 days of hospice care.8 The common missing element, whether patients die in the hospital or at home in hospice, likely remains a lack of early discussion and planning of care around their goals and priorities for the quality of the life they are leading as they experience serious illness.

Directions Forward
How is it possible to do better? It appears that the fundamental error has been to split the primary responsibilities for treatment of serious illness from providing care enabling the best possible quality of life in the face of such illness. Palliative care clinicians, geriatricians, and others have been doing essential work to develop, articulate, and deploy the skills required to serve the well-being of the sick and the frail. These patients have fears and worries about what is to come; they have goals and priorities for how they wish to live their lives; they have aspects of their lives they are willing to sacrifice, and aspects they are not willing to sacrifice, for the sake of more time; they have certain functions they consider essential for life to be worth preserving; and they want and need their caregivers and families to understand and support these concerns as a central part of their care.9

Such support is not a responsibility to be outsourced to a specialized few, however. For one, the supply is lacking—there are not nearly enough skilled palliative and geriatric specialists for the more than 2 million people per year in the United States who die from heart disease, cancer, chronic respiratory illness, cerebrovascular disease, Alzheimer disease, or end-stage renal disease, let alone around the world.2 Moreover, providing this support effectively requires integrating it into the care that everyone provides. There are reasonably well-established best practices for communicating prognosis, planning goals of care, and managing symptoms.9,10 However, training in using them has been absent from medical education in most specialties.

One view, often more implicit than explicit, is that the skills involved are not really that challenging—that clinicians are already perfectly capable of providing high-quality end-of-life care when they want to. Hence the popular claim that physicians receive different care when facing limited life expectancy—that physicians “die differently,” with more holistic, less heroic, and less institutional care than their patients receive. The implication is that physicians already know how to provide high-quality end-of-life care when they want to. Two

Research Letters in this issue of JAMA indicate that physicians do not die so differently, however.11,12 Both found that physicians were only a few percentage points less likely to die in the hospital than others, and Weissman et al found no difference in hospice use or total expenditures at the end of life. The hurdles are larger than mere motivation. Although the recently announced change in Medicare rules to provide payment for advance care planning conversations is welcome, necessary, and long overdue, evidence suggests it alone will not produce substantial improvement in patient care.

Physicians and other health professionals—even those with substantial experience caring for the seriously ill—commonly lack skills in eliciting the goals, preferences, and values of their patients and in effectively tuning their care to align with those aims. Incorporating these skills into practice will require dedicated education and coaching for trainees and practicing clinicians. It will also require their participation in research and innovation on how to best deploy their specialized capabilities at the end of life. Research in radiation oncology, for example, has established that single-fraction radiation therapy should be favored over multiple fraction therapy for palliation of bone metastasis.13 Likewise, Prigerson et al recently evaluated chemotherapy use in a prospective cohort of 384 patients with end-stage cancer (ie, progressive metastasis refractory to at least 1 line of chemotherapy) followed up until death.14 Half received palliative chemotherapy, but its use was not associated with improvement in either survival or quality of life in the last week of life, even when controlling for performance status. Indeed, patients with good baseline performance status—the group expected to achieve the most benefit—experienced a significantly worsened quality of life with chemotherapy use compared with none. This type of research, however, has been uncommon, particularly for non-oncologic conditions. Society has encouraged research talent and funding to focus narrowly on extending the quantity of life and not quality—another harmful consequence of pitting these 2 aims against one another.

Perhaps the most harmful splitting, however, has occurred in national policy concerning eligibility for hospice services. As Odejide15 notes in this issue of JAMA, Medicare and private insurers have codified benefit limits that require patients to give up on curative therapy in order to receive the intensive palliative services and management that hospice care provides. This has been a major contributor to the underuse of hospice, the underrecognition of the value of palliative care, and therefore the poor quality of life experienced by patients with advanced disease and limited life expectancy. This needs to change.

The recent Medicare demonstration project testing the lifting of those limits with 141 hospice organizations across the country is a welcome initiative.16 However, the evaluation runs the danger of continuing to treat support of quality of life differently from support of extension of life. Adoption as national policy should not depend merely on whether the change reduces overall health care costs. It may well do so, based on the experience of private insurers who have tested the idea. Nonetheless, just as with any other medical intervention, the key measure of effectiveness is how much it improves patients’ health and well-being.
The evidence indicates that the medical profession is harming vast numbers of patients by neglecting this goal—and that this is not just a US phenomenon but a global one. People everywhere have essential needs aside from just living longer. Medical practices, research, and policies must ensure that clinicians have the skills to understand those needs and have the capabilities to serve them for patients with life-limiting illness. Everyone dies. Death is not an inherent failure. Neglect, however, is.

Aims of Care
A bitter debate is continuing (including in this issue of JAMA)\(^\text{7,18}\) about the role of clinicians in enabling a good death for the small number of people who wish for assistance in dying. That discussion—about how best to balance respect for the sanctity of human life with the principle of autonomy—has drawn substantial public attention. However, the vital goal almost all people want from medicine is not having a good death but having as good a life as possible all the way to the very end.

REFERENCES
Sparrow FastCare and Sparrow Urgent Care offer the cure for the common wait with online check-in. Wondering if Sparrow FastCare, Sparrow Urgent Care, or the Emergency Room is best? Click here. Language Services. MyChart Bedside. With MyChart Bedside, Sparrow connects you to your care. Learn More. Click here. Lose Weight and Keep It Off. You can fit back into your life with a little help from Sparrow Bariatrics. Learn more by attending a free information session or clicking here. Healthy News. Sparrow Health System â€“ 1215 E Michigan Ave, Lansing, Michigan 48912 â€“ rated 4.1 based on 1,356 reviews "Visited FastCare in Frandor. I recommend signing...Â The report highlights how Sparrow is transforming care through our Caregivers, our national recognitions, the Sparrow Care Network, our Population Health initiative, award-winning specialties, and our Community Hospitals. The report also includes sections on the Sparrow Foundation, our finances, and governance bodies. The Annual Report website also contains a link to our new Nursing Annual Report. We hope the 2017 Sparrow Annual Report is a valuable resource about Sparrow and its 10,000 Physicians, Nurses, Volunteers, and other Caregivers. See more. See all.