

TO HELP YOU TO DIE WELL.





THE POWER OF STORY-WRITING THE BEST LAST CHAPTER

MY NAME IS MELODY. THIS IS THE STORY OF THE LAST CHAPTER OF THE LIFE OF MY HUSBAND



Dick wanted to be at home and I wanted him to be at home; to be together, and of course with our little dog. I wanted to be the one to care for him and he was comfortable with me looking after him. Some people are not comfortable with their spouse or family members performing some of the more personal and intimate care and prefer the more anonymous, professional, dispassionate assistance.

This is a very difficult time. This is a very personal decision. For us the decision was clear and obvious and we

were in complete agreement.

At home we were able (or not), bathing, watching TV, going to his computer, everything. Even visiting with neighbours and friends. That he /we made the right choice for us was confirmed for us. When he came home, we expected he'd have a week to 10 days and it turned out to be just a few days short of a year. Dick said he really believed that he would have been dead if he had to stay in the hospital. But more important than the length of time was the quality of time. The decline was inescapable, but Dick was LIVING HIS life right to the end.

Dick felt validated in having his decisions for

the end of his life just as he had for the rest of

his life. We were completely open and honest with each other, our family and friends, so everyone understood what he wanted and was

on the same page. That was important for me

him. He even went with me to the funeral home.

because everyone supported me supporting

to do everything on his time: sleeping, eating, dressing

We just said: "If the blinds are up, then we're up. C'mon in, and if he's up for a visit, then all's good. If it's not a good time, we'll tell you and you can try again later." There was no arbitrary schedule to be adhered to.

Our PHYSICIAN was available "whenever!" **HOME CARE** (we came in to the basement of the hospital) with bandages and butterflies as well as supplies, walker, lift chair, lift belt. **NUTRITIONIST** suggestions for

food choices and preparation.

PHARMACY kept ahead of the meds and sometimes sent them home with neighbours.

NURSE LINE (24 hours) which I called at 2:30 one morning for recommendation on pain meds. (She called back personally the next day to check how we were doing.)

The AMBULANCE / EMERGENCY room is always available.

Not under the umbrella of the medical community, but just as important was the support and help of our community, our friends and neighbours.

Palliative still means medical care, but being comfort, not cure. It was more important to Dick that he remained "himself" completely pain free, and I was attuned to signals and familiar Care does not come lived, but we had lots medical community.

The end was the next step along the same road, and felt like the natural transition. The experience was very personal and intimate. It was very hard for me but very rewarding. I knew that I had done the best that I could and that I had "done right by him". There was nothing left undone or unsaid. I have not had the experience of wishing for "just one more day". We were together, he was not alone, and it was clear that his life had run out naturally and completely. My grief is intense, but it feels clean and pure; it is not muddied with any guilt or regrets.

In the spirit of complete honesty, I will share that early on there were a couple of episodes of anger and frustration, on both of our parts. I don't know if it was his frustration with being sick or if he resented his declining abilities and having to give over what had always been his jobs to me. Maybe I just pissed him off. But I do know that passed and we grew even closer and more intimate at a deeper level than ever.

Every person's story matters. Loved ones need to be listened to and wishes need to be respected.





THE POWER OF STORY-WRITING THE BEST LAST CHAPTER

MY NAME IS PAT. THIS IS THE STORY OF THE LAST CHAPTER OF THE LIFE OF MY FATHER



Ken was born on December 12, 1933. He was a tremendous athlete growing up, but his love was hockey. He came to the East Kootenays in 1955 to play for the Trail Smoke Eaters. He moved to Kimberley the following season to play for the Dynamiters where he won several league and provincial hockey awards. After hockey, my dad stayed very active participating in golf, curling, cross country skiing and walking. He will also be forever remembered as the sales representative for Labatts

aka Kokanee during the 70s and 80s. Listen – KNOW WHO THE PATIENT IS - learn about their life, hopes, fears. Who are they, where have they been, what are they most proud of. What would you like your

caregivers to know about you?

Later in life, Ken faced several health challenges including bouts with esophageal and bladder cancer and developed poor heart health which led to having a pacemaker installed in 2016.

The beginning of the end began in the summer of 2017 when we noticed a definite loss of mobility and balance which eventually led to several falls in his home and while out walking. His partner, whose health was deteriorating from trying to take care of him, was forced to call an ambulance after a fall from which she could not help him up. He was taken to hospital.

It was suggested that I take him to my home which is not

set up for anything like that. So I arranged for him to move into a semi-assisted care facility. After one night there, he collapsed again and was taken back to the hospital by ambulance.

Information on community paramedic program; planning for long term care. WHO CAN HELP who needs to be included on the collaborative - integrated team?

Over the next week or so Dad was moved almost daily between the emergency room, day surgery room and the 2nd floor. I would visit my father every day but had no answers as to what was actually wrong with him. He continued to deteriorate and now his speech was becoming confused and unintelligible at times.

been helpful at this stage, allowing the family to prepare and identify their wishes. Introduce the palliative process in the latter part of the curative stage. ASK QUESTIONS - Who would you like to make decisions on your behalf if you can't?

Beginning palliative care would have

would be most make the right

On one visit the doctor wanted to meet with me to discuss my father. The floor was busy and the only place that was quiet was the ward TV room.

> There I was advised that my father was dying and that antibiotics were keeping him alive. The doctor wondered whether we should just keep him comfortable.

Disturbing discussion in a public venue. First time told Dad was dying. Possible explanation: "Your dad has had many illnesses and as we get older they take their toll, so he is dying from all those things."

Dying? That was a bit of a shock considering no one had ever, ever told me what was wrong with him. As I struggled to come to grips with what I was hearing, people were now moving in and out of the TV room. It was a very difficult situation I hadn't been given any information about Palliative Care.

After lengthy telephone conversations with my brother, we decided not to accept the doctor's recommendation and to carry on with treatment.

The conversations turned to what to do with my father. It was apparent the hospital wanted to discharge him as they felt that there was nothing more they could do for him. He could not go back to the semi-assisted care facility so I was again asked to move him to my home. If he was truly dying, my home and myself were even less prepared to take him than before. I was assured that home support would be provided but I felt that I was being given no options. I again refused.

need to be respected.

Finally, I was told that a respite bed had become available. On April 12, 2018 Dad took his last road trip in an ambulance to the care facility where he passed away on October 26, 2018.

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> > East Kootenay
> > Division of Family Practice



for them?