The use of induced pluripotent stem cells in domestic animals: a narrative review

Rachel A. Scarfone†, Samantha M. Pena†, Keith A. Russell, Dean H. Betts and Thomas G. Koch*

Abstract

Induced pluripotent stem cells (iPSCs) are undifferentiated stem cells characterized by the ability to differentiate into any cell type in the body. iPSCs are a relatively new and rapidly developing technology in many fields of biology, including developmental anatomy and physiology, pathology, and toxicology. These cells have great potential in research as they are self-renewing and pluripotent with minimal ethical concerns. Protocols for their production have been developed for many domestic animal species, which have since been used to further our knowledge in the progression and treatment of diseases. This research is valuable both for veterinary medicine as well as for the prospect of translation to human medicine. Safety, cost, and feasibility are potential barriers for this technology that must be considered before widespread clinical adoption. This review will analyze the literature pertaining to iPSCs derived from various domestic species with a focus on iPSC production and characterization, applications for tissue and disease research, and applications for disease treatment.

Keywords: Induced pluripotent stem cells, Domestic species, Veterinary medicine, Production, Characterization, Disease modelling, Disease treatment

Background

Induced pluripotent stem cells (iPSCs) are laboratory-developed pluripotent stem cells generated by the reprogramming of differentiated cells [1]. Takahashi and Yamanaka first discovered somatic cells’ capacity for reprogramming in 2006 after forcing differentiated fibroblast cells to ectopically express four transcription factors associated with pluripotency: Oct4, Sox2, Klf4, and c-Myc, collectively referred to as OSKM [1, 2]. iPSCs have since been of interest to researchers in the fields of toxicology, pathology, virology, developmental anatomy and physiology, amongst others [3–5]. iPSCs possess several benefits over other stem cell types such as mesenchymal stromal cells (MSCs) and embryonic stem cells (ESCs). In the context of this review, the term mesenchymal stem cells is preferred over mesenchymal stem cells due to the finite self-renewing property of MSCs that does not support the traditionally recognized self-renewing characteristic of stem cells [6]. The versatility of iPSCs may make them preferential over MSCs that are limited in their differentiation potential due to their multipotent nature [7–9]. ESCs offer a similar versatility to iPSCs as they are both pluripotent, but not without limitations [8]. ESCs can be obtained from in vivo and in vitro produced embryos at the blastocyst stage [10]. However, technical difficulties have interfered with the isolation and use of ESCs, namely in ungulate species and canines [2, 8, 11, 12]. Oocyte collection for in vitro embryo production is an invasive procedure that has prompted ethical considerations. Disposed reproductive material has been the primary source of oocytes in domestic species obtained from meat processing in livestock or ovariohysterectomies in companion animals [13–15]. In vivo protocols may include minimally invasive uterine flushing, often seen in mares [10]. iPSCs provide a more practical alternative to...
creating ESC-like cells in species where recovery of embryos or in vitro fertilization is difficult or not possible [12]. Unlike ESC lines, autologous iPSC lines can also be produced. This is ideal for transplantation of stem cells and their derivatives as it avoids the immunological complications associated with allogeneic iPSCs. Consequently, iPSCs can be used as an alternative to MSCs and ESCs with the potential for greater research and clinical applicability in domestic species.

While research has focused primarily on human and mice iPSCs, there has been a slow accumulation of iPSC research in domestic animals in the last decade (Fig. 1). iPSC derivation protocols have been developed in species including porcine [16], equine [17], canine [18], bovine [19], galline [20], caprine [21], ovine [22], and feline [23]. Aside from their importance in treating veterinary pathologies, porcine, canine, and equine models have been shown to be valuable for the study and treatment of human disease [24–26]. The purpose of this review is to provide an overview of the literature pertaining to current protocols and applications of iPSCs derived from domestic species. This review will address the topics of the development and use of iPSCs for tissue and disease research, their treatment in domestic animals and the barriers to their production and applications.

**iPSC production and characterization**

Yamanaka and colleagues’ discovery of iPSCs originated in mice models, followed closely by their derivation from human fibroblasts [1, 27]. Briefly, mice tail fibroblasts or human dermal fibroblasts were cultured then transduced with retroviral vectors containing expression cassettes of the OSKM reprogramming factors, inducing pluripotency in the transduced cells (Fig. 2). Using these protocols as a base, methods have been adapted in order to produce iPSCs in other species.

iPSCs have been developed from porcine [16], equine [17], canine [18], bovine [19], galline [20], caprine [21], ovine [22], and feline [23] tissue. Successful iPSC production from domestic species was first reported in 2009 by Wu and colleagues in porcine, and the field has since expanded to other species (Fig. 1). iPSCs have been produced from various donor tissue types, transduction systems, and reprogramming factor combinations. In domestic species, iPSCs have been derived from fibroblasts, MSCs and other somatic cell types including epithelial and testicular cells (Table 1). Tissue sources have been obtained from various developmental stages, namely fetal, neonatal, juvenile, and adult. For simplicity, this review has identified any tissue sources obtained from an animal in utero as fetal and those obtained after birth as adult. Deriving iPSCs from adult somatic cells is generally preferable to embryonic derivation due to a higher abundance of cells, easier collection of cells, and the ability to produce autologous iPSC populations for disease treatment. Donor tissue is then cultured and reprogrammed using viral or non-viral vectors containing the designated reprogramming factors. Viral vectors include lentiviruses, oncoviruses, and Sendai viruses, while non-viral vectors include cDNA vectors, minicircles and transposons (Table 1). The selected reprogramming factors typically include OSKM, but other variations have also been explored. Nanog and Lin28 are commonly used in the literature in addition to OSKM, and a small number of papers report the use of other additional transcription factors, such as TERT, and Tet1 (Table 1). More recently, work has been carried out using microRNAs in combination with other factors to achieve pluripotency induction [19, 29, 67]. MicroRNAs alone have only shown partial reprogramming abilities in domestic animals [61].

![](cumulative_iPSC-related_publications.png)

**Fig. 1** Cumulative iPSC-Related Publications in Domestic Species, January 2008–March 2020. **A** Publications regarding induced pluripotent stem cells from January 2008 to March 2020 in domestic animal species including porcine, equine, canine, bovine, galline, caprine, ovine, and feline. Increased interest in iPSC research in domestic animals is demonstrated, particularly in the porcine model. **B** A subset of publications excluding porcine papers to visualize the general positive trend in all other domestic species.
Following presumed iPSC production, colonies can be analyzed via morphological assessment to select for colonies with the most potential in reprogramming cells to an undifferentiated state. Non-invasive morphological assessment also provides insight into the developmental competence and homogeneity of iPSC colonies. Traditionally, iPSC colonies resemble ESC colonies with well-defined borders and tightly packed cells. More specifically, dome-shaped and flattened colonies are indicators of naïve and primed pluripotency, respectively [111].

Cells in these colonies are expected to have a large nucleus and little cytoplasm [112]. Naïve pluripotency is recognized by characteristic molecular features of the pre-implantation mouse embryonic stem cell, whereas primed pluripotency resembles stem cells of the post-implantation mouse epiblast [113]. Naïve pluripotent stem cells are identified by X chromosome reactivation in females, dependency on leukemia inhibitory factor (LIF) and receptivity to BMP4 to maintain pluripotency, and the transition to a more differentiated state in response to FGF2 and ACTIN/TGFB signalling [58, 114, 115].

Putative iPSCs must then undergo a series of tests to confirm pluripotency (Table 2). In domestic species, pluripotency is often confirmed by the endogenous expression of pluripotency markers, and the formation of in vitro embryoid bodies and in vivo teratomas containing cell types derived from all three germ layers [118]. Chimera formation with germ-line transmission is a less commonly used method in domestic species (as demonstrated in Table 2), but is deemed the gold standard for validating stem cell pluripotency [119].

Although formation of embryoid bodies and teratomas are successful in the majority of papers referenced in this review, many publications lack complete pluripotent characterization of their produced cell lines. Consequently, this review uses the term iPSCs broadly to describe both bona fide iPSCs and iPSC-like cells as some primary research lacks sufficient iPSC characterization.
| Origin cell type                        | Reprogramming system               | Reprogramming factors                        | Reference |
|----------------------------------------|------------------------------------|---------------------------------------------|-----------|
| **Porcine**                            |                                    |                                             |           |
| Fetal fibroblasts                      | Unspecified retroviral vectors     | OSKM                                        | [29–38]   |
| Fetal fibroblasts                      | Oncoviral vectors                  | OSKM                                        | [39]      |
| Fetal fibroblasts                      | Lentiviral vectors                 | OSKM                                        | [16, 40–46]|
| Adult sertoli cells                    | Unspecified retroviral vectors     | OSKM                                        | [47, 48]  |
| Adult fibroblasts                      | Unspecified retroviral vectors     | OSKM                                        | [49]      |
| Adult fibroblasts                      | Lentiviral vectors                 | OSKM                                        | [24, 50–53]|
| Adult fibroblasts                      | Sendai viral vectors               | OSKM                                        | [54]      |
| Adult fibroblasts                      | Lentiviral vectors                 | OSKM, Nanog, Lin28                          | [55, 56]  |
| Adult MSCs                             | Lentiviral vectors                 | OSKM, Nanog, Lin28                          | [57]      |
| Adult MSCs and fibroblasts             | Lentiviral vectors                 | OSKM                                        | [58]      |
| Adult fibroblasts and bone marrow cells| Lentiviral vectors                 | OSKM, Nanog, LIN28                          | [59]      |
| Fetal fibroblasts                      | PiggyBac transposon               | OSKM                                        | [60]      |
| Fetal fibroblasts                      | Lentiviral vectors                 | miR-302 s                                   | [61]      |
| Fetal fibroblasts                      | Unspecified retroviral vectors     | OSKM                                        | [62]      |
| Fetal fibroblasts                      | Episomal plasmids                 | Oct3/4, Sox2, Klf4, I-Myc                    | [63]      |
| Adult fibroblasts                      | Lentiviral vectors                 | OSKM, Nanog, Lin28                          | [64]      |
| Fetal fibroblasts                      | Unspecified retroviral vectors     | OSKM, mTet3, Tet1, Kdm3a                     | [65]      |
| Fetal fibroblasts                      | Lentiviral vectors                 | OSKM, Nanog, Lin28                          | [66]      |
| Fetal fibroblasts                      | Unspecified retroviral vectors     | OSKM, miR-106a-363, and miR-302             | [67]      |
| Fetal fibroblasts                      | Lentiviral vectors                 | OSKM, or OSKM, Tbx3, Nr5a2                  | [68]      |
| Fetal fibroblasts                      | Sleeping Beauty transposon         | OSKM, Nanog, Lin28                          | [69]      |
| Fetal fibroblasts                      | Unspecified retroviral vectors     | OSKM, TERT                                   | [70]      |
| Fetal and adult fibroblasts and MSCs   | Unspecified retroviral vectors and lentiviral vectors | OSKM | [71] |
| Adult fibroblasts                      | Unspecified retroviral vectors     | OSKM                                        | [72]      |
| Fetal fibroblasts                      | Sleeping Beauty transposon         | OSKM, Nanog, Lin28                          | [73]      |
| Fibroblasts                            | Sleeping Beauty transposon         | OSKM                                        | [74]      |
| GALT-KO fibroblasts                    | Lentiviral vectors                 | OSKM, Nanog, LIN28                          | [75]      |
| Adult fibroblasts                      | Lentiviral vectors                 | OSKM, Nanog, Lin28                          | [76]      |
| Adult MSCs                             | Lentiviral vectors                 | OSKM                                        | [77]      |
| Fetal fibroblasts                      | Episomal vectors                   | Oct4, Sox2, Klf4                            | [78]      |
| Fetal MSCs                             | Unspecified retroviral vectors     | Oct4, Klf4                                   | [79]      |
| Fetal fibroblasts                      | Lentiviral vectors                 | OSKM, Nanog                                 | [80]      |
| Adult fibroblasts                      | Unspecified retroviral vectors     | OSKM                                        | [81]      |
| **Equine**                             |                                    |                                             |           |
| Fetal fibroblasts                      | PiggyBac transposon               | OSKM                                        | [17]      |
| Adult fibroblasts                      | Unspecified retroviral vectors     | Oct4, Sox2, Klf4                            | [82]      |
| Adult fibroblasts                      | Unspecified retroviral vectors     | OSKM                                        | [83]      |
| Adult fibroblasts                      | PiggyBac transposon               | OSKM                                        | [84, 85]  |
| Adult keratinocytes                    | Unspecified retroviral vectors     | OSKM                                        | [86]      |
| Adult MSCs                             | Lentiviral vectors                 | OSKM                                        | [87]      |
| **Canine**                             |                                    |                                             |           |
| Fetal fibroblasts                      | Unspecified retroviral vectors     | OSKM                                        | [18]      |
| Fetal fibroblasts                      | Sendai viral vectors               | OSKM                                        | [88]      |
to confirm true pluripotency. A common occurrence among studies is the incomplete silencing of exogenous transcription factors. These cell lines may not be truly pluripotent, but are reliant on transgenes to maintain pluripotency. The use of epigenetic modifiers has previously been shown in human and mice cells to increase transgene silencing while maintaining endogenous pluripotency factor expression; results that have now been replicated in the porcine model [65]. This technique could be promising to alleviate these issues, although more work will be required.

As previously mentioned, the gold standard for validating stem cell pluripotency is via chimera formation with germ-line transmission. Chimera formation is defined by the heterogeneous cell population of an early embryo following the injection of iPSCs into the blastocyst. This confirmation method requires iPSCs to integrate into the developing embryo and contribute to all three germ layers and, potentially, germ cells. To confirm germ-line transmission, chimeras are mated with non-chimeras and offspring are assessed for iPSC contribution [116]. Previous research has suggested that the
feasibility of chimerism and germ-line transmission is greatly improved with the use of naïve pluripotent stem cells, as opposed to primed pluripotent stem cells [2, 40]. Although porcine naïve iPSC-like cells have been reported, there is little evidence of germline transmission. Analysis of the literature suggested that the generation of chimeras in livestock species is difficult to achieve [20]. Injection of generated iPSCs into an embryo often resulted in limited incorporation [40, 108] and the resulting offspring often were not chimeric [40, 108, 117]. Significant variation in iPSC integration has been shown [19, 112]. To the best of our knowledge, West and colleagues remain the only researchers to successfully realize germline transmission of iPSCs in a domestic species (i.e. porcine) [116]. iPSCs were also tested in cloning transgenic tissue [55, 104] and genomic incorporation of transgenes [60]. It was found that iPSCs could be derived from transgenic organisms, specifically genetically modified pigs designed for xenotransplantation [55]. However, limited developmental potential of embryos past the blastocyst stage was observed [60]. Hence, analysis of germline transmission is currently not feasible in such species. In moving forward with clinical applications of iPSCs in either human or veterinary medicine, being able to truly define cells as iPSCs will be crucial for standardization, quality, and safety assurances.

**Tissue and disease research**

iPSCs have the potential to be valuable tools for tissue and disease modelling. In vitro differentiation of iPSCs has allowed for study of the developmental processes and pathologies of tissues and may allow for preclinical testing of therapeutic drugs for veterinary and human medicine. With regard to drug screening, there has been success in human and mouse iPSC research in using differentiated iPSC lines to model disease and conduct high-throughput screening of small molecules for their effects on disease progression [120]. This technique allows for testing of potential therapeutics against disease-genotype cells specific to an individual or species without the need for interspecies comparisons or excessive lab animal use. Differentiation into specific cell types has been noted many times in the literature in porcine, equine, canine, galline, and bovine models, which are described below. Although characterization of these differentiation cells is demonstrated by physiological, genetic, or metabolic capacities of cell lines, the degree of differentiation varies from progenitor cells (e.g. neural progenitors [41, 76, 121]), to fully differentiated cell types (e.g. skeletal myocytes [122]). Domestic animal diseases are abundant and have negative health effects for consumers of agricultural animal by-products [123–126]. Unfortunately, the use of stem cells for research on livestock disease is novel and presently limited in number. The prolonged self-renewing characteristic of iPSCs supports their use in the study of physiology, disease pathology, drug toxicity and vaccine development in domestic species. A summary of veterinary animal iPSC research can be found in Table 3.

**Porcine**

Porcine iPSCs (piPSCs) have been differentiated into several cell types for research purposes. Currently, they have been used in the production of neural progenitor cells [41, 76, 121], endothelial cells [30], myotubes [134], hepatocytes [50, 78], and vascular smooth muscle cells (VSMCs) [42]. VSMCs in particular have been applied to scaffolds for implantation into immunodeficient mice and successfully formed 3D scaffold-free tissue rings [42].

| Origin cell type | Suggested pluripotent state | iPSC characterization criteria | Reference |
|------------------|-----------------------------|--------------------------------|-----------|
| **Porcine**      |                             | Pluripotency markers Embryoid bodies Teratomas Chimeras Germline transmission |           |
| Adult MSCs       | Primed                      | Yes                           | Yes       | Not tested | Yes       | Yes       | [57, 116] |
| Fetal fibroblasts| Naïve                       | Yes                           | Yes       | Yes; limited to blastocyst | Not tested | [117]     |
| Fetal fibroblasts| Naïve                       | Yes                           | Yes       | No         | Yes; limited to fetus | Not tested | [40]      |
| Fetal fibroblasts| Primed                      | Yes                           | Yes       | Yes; but low contribution at birth | Not tested | [108]     |
| **Ovine**        |                             |                                |           |            |           |           |           |
| Ovine fibroblasts| Primed                      | Yes                           | Yes       |            |           |           |           |
| **Galline**      |                             |                                |           |            |           |           |           |
### Table 3 iPSC publications relating to tissue and disease modelling

| Origin cell type | Differentiated cell type | Tissue or disease target | Outcome | Reference |
|------------------|--------------------------|--------------------------|---------|-----------|
| **Porcine**      |                          |                          |         |           |
| Fibroblasts      | Neural Rosettes, neural crest-like cells, and peripheral sensory neural-like cells | Neural tissue | Indications of neural differentiation by the upregulation of sensory neuron genes and peripheral neuron markers | [121] |
| Adult fibroblasts| Neurons, astrocytes, and oligodendrocytes | Neural tissue | Indications of neural differentiation by the presence of mature neural markers and morphology of neurons, astrocytes, and oligodendrocytes; indications of further differentiation into motor neurons. | [76] |
| Fetal fibroblasts| Neural progenitor cells | Neural tissue | Production of neural progenitor cells with expression of neuronal markers | [41] |
| Fetal fibroblasts| Endothelial cells | Endothelial tissue | Production of endothelial cells with morphological and functional properties | [30] |
| Fetal fibroblasts| Hepatocyte-like cells | Liver tissue | Production of differentiated cells characteristic of hepatocytes by functional properties | [78] |
| Adult fibroblasts| Hepatocyte-like cells | Liver tissue | Production of differentiated cells characteristic of hepatocytes functional properties | [50] |
| Fetal fibroblasts| Vascular smooth muscle cells | Muscle tissue | Production of vascular smooth muscle cells capable of forming 3D scaffold-free tissue rings | [42] |
| **Equine**       |                          |                          |         |           |
| Adult fibroblasts| Cortical neurons | West Nile Virus (WNV) and Murray Valley Encephalitis (MVEV) | Successful infection of functional eiPSC-derived neurons by WNV and MVEV | [127] |
| Adult keratinocytes| Cholinergic motor neurons | Motor Neurons | Production of functional neurons capable of generating action potentials | [86] |
| Fetal and adult fibroblasts| Tenocytes | Tendons | Formation of three-dimensional artificial tendons | [128] |
| Fetal fibroblasts| Skeletal myotubes | Muscle Tissue | Formation of eiPSC-derived muscle fibers with electrophysiological function | [122] |
| Adult fibroblasts| Osteoblasts | Bone | Formation of eiPSC-derived bone tissue capable of secreting hydroxyapatite and calcium matrix | [129] |
| Adult keratinocytes| Primary keratinocytes | Epidermal wounds | Creation of artificial tissues for potential skin graft applications | [84] |
| **Canine**       |                          |                          |         |           |
| Adult fibroblasts| MSCs | Cartilage and Bone Tissue | Formation of three-dimensional chondrogenic and osteogenic cultures | [130] |
| Fetal fibroblasts| Mature megakaryocytes | Thrombocytopenia | Production of cells capable of releasing functional platelets upon signaling induction | [131] |
| **Bovine**       |                          |                          |         |           |
| Adult testicular cells| N/A | Phthalate ester exposure | Significant reduction in androgen expression and increase in apoptosis | [94] |
| Adult epithelial cells| Mammary epithelial-like cells | Mammary tissue | Indication of mammary phenotype for iPSCs cultured with progesterone | [97] |
| **Galline**      |                          |                          |         |           |
| Fetal fibroblasts| N/A | Goose influenza H5 | Incorporation of replication-incompetent virus into iPSCs | [100] |
| Fetal fibroblasts| N/A | Newcastle disease virus (NDV) | Successful infection of iPSCs with NDV; viable iPSCs exhibited increased tolerability | [132, 133] |
Equine

Equine iPSCs (ePSCs) have been differentiated into several cell and tissue types for disease modelling including neurons [86, 127], tendons [128], myotubes [122], and osteoblasts [129]. Functional ePSC-derived neurons have been produced and were capable of firing action potentials in vitro via functional calcium channels [86]. One paper reported the observation of neurospheres with axonal outgrowths connecting adjacent cells [127]. This paper studied the potential for neurospheres to model West Nile virus (WNV) and Murray Valley encephalitis virus (MVEV), infectious, neurotropic equine diseases [127]. iPSC-derived neurons were successfully infected by WNV and MVEV, which could allow for future research to study mechanisms of these and other infectious diseases and neuropathic conditions.

Musculoskeletal tissue is a major system that would benefit from ePSC modelling due to the frequency of injuries in competing horses. Artificial tendons derived from iPSCs have been attempted and although two-dimensional assays showed matrix contraction and appropriate gene expression, three-dimensional assays failed to generate functional artificial tendons. ESCs were shown to more efficiently produce functional tendons [128]. Further study is required here as this could be a promising area of regenerative medicine if ePSC-derived tendons can be improved. Using fibroblast-derived ePSC lines, researchers induced differentiation into myocytes, the functional unit of muscles. Myotubes demonstrated intracellular calcium release following membrane depolarization [122]. Lastly, functional ePSC-derived osteoblasts have been reported. These cells expressed genetic markers of osteoblasts and were shown to produce hydroxyapatite and calcium matrices, highly specific characteristics of bone tissue [129]. Artificial production of bone may allow for study of bone physiology and diseases but may also benefit veterinary treatment of fractures and other pathologies.

Wound management is a common problem in equine medicine, and skin grafting, the ideal treatment, is often not possible due to a low supply of donor tissues [135]. One paper described a protocol where ePSCs were differentiated into keratinocytes (ePSC-KCs) to produce skin grafts. The ePSC-KCs were likened to both progenitor and primary keratinocyte-like cells, potentially indicative of epidermal basal stem cell identity, ideal for in vivo wound management [84].

Canine

MSCs derived from canine iPSCs (ciPSCs) have been proposed as an intermediate stage to developing canine models of musculoskeletal tissues through chondrogenic and osteogenic pathways [130]. ciPSCs were differentiated into MSCs and subsequently differentiated into chondrocytes and osteoblasts in three-dimensional hydrogel culture conditions. Researchers proposed these three-dimensional cultures as effective models for studying canine osteoarthritis in order to develop MSC-based therapies and further model human degenerative joint disease [130].

A novel protocol has been published to generate functional canine platelets to treat thrombocytopenia, a canine and human clotting disorder. ciPSCs were differentiated into mature megakaryocytes which could be induced to release functional platelets [131]. This could serve as an alternative treatment to blood transfusion, the only effective therapy currently available.

Galline

Galline iPSCs (giPSCs) have been used in studying viral infection and replication [100, 132, 133]. Newcastle disease (NDV) is a common avian viral disease often found in domestic poultry [132]. Studies have demonstrated that giPSCs are capable of NDV infection [132, 133], and that viable cells displayed increased tolerability but not immunity to the virus [133]. giPSCs could also be used to produce replication-incompetent viruses, such as the highly pathogenic H5 avian influenza viruses [136]. Replication-incompetent viruses were produced with the goose influenza H5 gene and were incorporated into giPSCs. Using these cells, the virus was further transduced into a bladder cancer-derived cell line and could be inactivated by formaldehyde [100]. The use of giPSCs for vaccine production may be beneficial over chick embryos or eggs due to a decreased risk of contamination [100]. These results suggest that giPSCs have the potential to produce inactive viruses for vaccine production.

Bovine

Limited disease modelling has been observed with bovine iPSCs (biPSCs); however, current efforts have demonstrated their potential application in toxicological studies to elucidate the effects of toxic environmental compounds. Cattle can be used to investigate the negative effects of environmental endocrine disrupting compounds (EDCs) on humans and livestock species as the potential for harmful chemicals leaching into waterways and soils has become a prominent concern [137]. Despite a lack of clinical evidence, it has been proposed that EDCs can affect the reproductive functioning of cattle, greatly impacting agricultural production [138]. Bovine iPSCs have been applied to research the EDCs phthalate esters [94]. It was found that phthalate esters significantly downregulated androgen receptors of iPSCs, which supported apoptosis [94]. Such studies introduced biPSCs as a feasible tool in studying the effects of endocrine disruptors and other chemicals on cell populations.
bipSCs have also been differentiated into epithelial-like cells that phenotypically resembled mammary cells [97]. These cells could further be investigated for their application in tissue regeneration for oncology patients who have undergone a mastectomy.

Disease treatment
Although research of specific pathologies is generally limited to single publications, the use of iPSCs to treat diseases and injuries in animals is growing and will likely be integrated into veterinary practice in the future. The field of domestic animal regenerative medicine may also provide models for human pathologies. Stem cell research that was once conducted on rodents is now growing in dogs and pigs [139, 140], species shown to be better models for human disease [25, 139, 140]. Table 4 summarizes the current research for iPSC-based treatments in domestic animals, which for the purpose of this review, includes all in vivo applications of iPSCs and their derivatives.

Porcine
Pigs are the most frequently used model of disease in domestic species. Porcine iPSCs have been employed in the study of tissue regeneration in bone [24, 141], muscle [51, 142], and nervous tissue [54, 143]. The findings in a majority of the articles published confirms that piPSCs are capable of integrating into tissue at the site of implantation [142, 143] and are capable of cueing endogenous pathways to upregulate [51], thus improving conditions at the site of tissue damage or death.

In a study of bone regeneration, piPSC-derived osteoblast-like cells were able to improve the trabecular and cortical bone structures of fractured tibias [141]. In a similar study, partial tibial cartilage regeneration at the transplantation site was observed with the regenerated cartilage originating from iPSCs [24].

Table 4 iPSCs for Disease Research

| Disease Target                  | Origin Cell Type | Differentiated Cell Type | Route of Administration | Outcome                                                                                                                                                                                                 | Reference |
|---------------------------------|-----------------|--------------------------|--------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------|
| Porcine Osteoporosis            | Fibroblasts     | Osteoblast-like cells    | Local cell transplantation | Significant improvement in bone structures at transplanted site; maintenance of bone structures locally                                                                                                    | [141]     |
| Porcine Osteochondral damage, osteoarthritis | Adult fibroblasts | piPSC-like cells         | Direct pellet transplantation | Cartilage regeneration; no tumor formation                                                                                                                                                               | [24]      |
| Porcine Chronic myocardial infarction | Adult fibroblasts | piPSCs                  | Direct injection          | Integration of iPSCs into cardiac muscle without differentiation; potential contribution to angiogenesis                                                                                             | [51]      |
| Porcine Acute myocardial infarction | Adult fibroblasts | piPSCs                  | Direct injection          | Significant decrease in infarcted area; improvement in local function and perfusion                                                                                                                    | [142]     |
| Porcine Myocardial infarction    | Adult MSCs      | Endothelial cells (ECs)  | Local injections          | Improved function and an increase in the number of capillaries in the peri-infarct area; no significant changes in infarct area size.                                                                  | [77]      |
| Porcine Chronic spinal cord injury | Adult fibroblasts | Neural precursor cells (NPCs) | Bilateral syngeneic grafts | Long-term immune tolerance of NPCs; integration into and beyond grafted region                                                                                                                         | [54]      |
| Porcine Retinal damage          | Fetal fibroblasts | Rod photoreceptors       | Local injection           | Integration into damaged porcine neural retina                                                                                                                                                           | [78, 143] |
| Equine Muscle injury            | Adult MSCs      | eIPSCs                   | Intramuscular injection   | Partial muscle regeneration; in vivo differentiation of eIPSCs into myofibers at the injury site                                                                                                           | [87]      |
| Equine Musculoskeletal injury    | Adult MSCs      | MSCs                     | Injection into lesion     | Improvements in clinical conditions for injuries including fractures, tendinitis, osteochondrosis, and osteoarthritis                                                                                   | [144]     |
| Canine Hind limb ischemia       | Adult MSCs and fibroblasts | Endothelial cells        | Local injections          | Successful engraftment in the ischemic limb; significant improvement of vascularization locally                                                                                                       | [92]      |
| Canine Cardiac infarction       | Adult MSCs and fibroblasts | Endothelial cells        | Local injections          | Successful engraftment locally; improvement in cardiac contractility                                                                                                                               | [92]      |
Other studies examined the beneficial treatment effect of iPSCs on chronic myocardial ischemia [51] and infarction [142]. Regenerative therapy of cardiac tissue in porcine models involved direct injection of undifferentiated piPSCs into myocardium [51, 142]. The treatment was found to significantly decrease the infarction area, decrease regional perfusion, and increase angiogenesis with local incorporation of piPSCs into myocardium and blood vessel without tumor formation [142]. A similar study found small tumor formations that eventually arrested in growth [51]. While the research suggested variations in the grafting capabilities of these piPSCs into host tissues, there was an identified increase in smooth muscle actin, indicating piPSCs interact in some form with host tissue. In brief, piPSCs have been shown to contribute to myocardium regeneration.

piPSCs were also applied to regenerate nerve tissue. Researchers differentiated porcine iPSCs into neural progenitor cells (NPCs) and rod photoreceptors in vitro, then successfully implanted them into the site of cell damage. piPSCs not only incorporated into the host tissue at the site of implantation, but further extended beyond the grafted region long-term [54, 143]. The results suggest that piPSCs are capable of effectively integrating into host tissue, making them a candidate for clinical application.

Equine
Two papers have been published describing an in vivo application of eiPSCs for the treatment of musculoskeletal injuries in equines [87, 144]. In the first paper, published in 2016, muscle injuries were induced in a GFP mouse model by injecting notexin, a myotoxic venom, along with an injection of eiPSCs. It was reported that these muscles saw an increase in myofiber production, and since eiPSCs were non-GFP reporting, it was shown that muscle fibers originating from the eiPSCs were produced. Undifferentiated cells remained in the muscle, indicating the dangerous potential for cancer formation [87].

To safeguard against potential cancer formation, another paper differentiated eiPSCs into MSCs prior to injection, reducing the risk of undesired proliferation. The eiPSC-MSCs were then injected into horses with various musculoskeletal disorders including fractures, tendonitis, osteochondrosis, and osteoarthritis. Improvements were observed including reduced lameness fever and fracture lines, although some horses also experienced hot flush and edema [144]. Although successful, this paper indicates a need for further development of less immune-reactive therapies.

Host immune responses are a major concern for clinical use of iPSCs, especially in species like horses where allogeneic cell use would be ideal. Further to the example above, another paper tested the immune potential of in vivo transplantation of allogeneic eiPSCs. Injected cells induced a minor, focal inflammatory response, but cellular signs of chronic inflammation persisted until the end of the study period 30 days after grafting. Undifferentiated cells have reduced expression of MHC surface proteins, but upon differentiation in vivo, these proteins increase, stimulating an increased immune reaction [145]. Although these cells were undifferentiated, the risk of immune response is significant and must still be addressed in differentiated eiPSCs, especially if this response increases with differentiation prior to implantation.

Canine
Fewer developments have been made in ciPSC research, but a 2011 paper showed the potential for ciPSCs to be used for ischemic tissue damage treatment, both in hind limb ischemia and cardiac infarction mouse models [92]. ciPSCs were differentiated into endothelial cells (ciPSC-ECs), then injected into mice models. In hindlimb ischemia mice, ciPSC-ECs were shown to significantly improve revascularization in the compromised tissue. In cardiac infarction models, ciPSC-ECs were shown to engraft onto the heart muscle itself and improve cardiac contractility. In both models it was demonstrated that donor cells were lost over time, indicating a possible need for repeated treatments. Nevertheless, the lack of recurring original symptoms of ischemia suggests the capability of these cells to induce long-lasting, persistent effects in tissues following their disappearance [92].

Barriers
Safety
There are several concerns to be resolved before in vivo use of iPSCs can be justified. Immune reactivity is one concern in the use of allogeneic cells that has been discussed earlier in this view. Aside from the formation of undesired cell types, the most significant risk is in vivo tumorigenesis due to the proliferative potential of iPSCs. Current research has demonstrated that differentiation of iPSCs and purification of differentiated cellular products prior to implantation can reduce tumour formation [146]. Alternatively, tumour formation has been addressed in mice models with the application of “suicide genes”. Using a drug-inducible suicide system, apoptosis of iPSC-derived cells can be initiated with exposure to a particular drug [147]. This system would allow for the complete inactivation of iPSC derivatives in the event of aberrant growth or modification.

Immune reaction to transplanted iPSCs is another safety concern for clinical application. The use of autologous transplants would mitigate these effects, although is not realistic for commercialization of treatments due to prohibitively high costs. Research is being conducted
into a cellular “cloaking” system that would allow cells to go undetected by the immune system of the host. Modification of allogeneic cells by altering MHC and HLA antigens has also shown potential to eliminate immune reaction in human iPSCs xenotransplantation studies [148]. However, this system is not without limitations; it is dangerous to create cell populations that cannot be controlled by the host immune system. Introducing a system that combines the drug-inducible suicide system and the cloaking system could potentially resolve this issue.

Many transduction systems used for iPSC production have inherent safety concerns due to their random integration into the host genome. Random integration can lead to disruptions in host genes and an increased risk of oncogenicity. A non-genome integrating Sendai virus system [149, 150] allows for the production of transgene-free iPSCs while maintaining a high reprogramming efficiency [149]. Similarly, a non-viral system that operates on the use of piggyBac transposons can create transgene-free iPSCs via excision from the genome following iPSC generation [151]. Originally described in human models, these systems have since been applied to domestic species, including dogs [88, 89], chickens [20], and cattle [152]. The wide range of reprogramming system options is beyond the scope of this article but have been reviewed elsewhere [153–155].

Technical barriers
Researchers have relied on precedent methods of human and mouse models to generate iPSCs in domestic animals [49, 114, 156]. An issue seen in many domestic models is the retention of pluripotent transgene expression; a situation that allows for the maintenance of pluripotency, or in many cases a pseudo-pluripotent state, that can interfere with differentiation. Most iPSCs derived from domestic species have been generated by viral integration of human or murine reprogramming transgenes that remain expressed [82, 83, 85, 86, 91]. The continuous expression of these transgenes suggests an incomplete epigenetic remodeling with OKSM factors alone and a greater need for understanding and optimizing the pluripotency induction process in domestic species. The use of non-viral vectors may prove effective in iPSC production, while overcoming the issue of transgene expression. Unfortunately, there is limited research on the application of iPSCs for disease research with the use of non-viral vectors. Yu and colleagues remain the only research group to successfully generate iPSCs using non-viral minicircles capable of generating chimeric chicks [20]. As few researchers have confirmed pluripotency by means of chimerism, confirmation of bona fide iPSCs has been limited. Often, cells believed to be iPSCs are iPSC-like cells as there are technical difficulties in yielding bona fide iPSCs that can maintain pluripotency independent of doxycycline [80]. Bona fide iPSCs remain difficult to obtain and further investigation into true iPSC production is required.

Despite significant species conservation of pluripotency genes, some divergence of the core pluripotency genes have been identified between mammals [157]. For example, the use of OSKM, Lin-28 and Nanog has been well established in porcine models, while other species are still under investigation, e.g. felids where OSKM plus NANOG may be required [23, 158]. It may be necessary to modify existing methods of achieving pluripotency, such as including additional reprogramming factors or developing different culture conditions to overcome species-specific reprogramming barriers.

Cost
Cost is another barrier to the application of iPSCs in domestic species due to laborious production. As previously mentioned, iPSCs from domestic animals have technical barriers limiting yield. As a result, more reagents, tissue, time, and labour are required for sufficient production [159]. Further costs have been associated with autologous iPSC treatment due to the production and maintenance of many cell lines and associated labour costs.

Future directions
Organoids, three-dimensional cell cultures that demonstrate characteristic development, anatomy, and physiology of a tissue, are currently an undeveloped tool in iPSCs derived from domestic species. Organoids have recently been developed from human iPSCs, which suggests the possibility of producing any organ of the body under the appropriate conditions. The use of iPSCs derived from domestic animals for organoid production could also be applied to veterinary medicine. Similar to human iPSCs, two-dimensional models have limitations in drug screen and assessing disease progression as they do not resemble in vivo conditions like organoids. Hence, virologists and drug developers can use them to better understand the mechanisms of disease or drug actions [160]. Several researchers have already exemplified the use of human embryonic stem cell-derived organoids in detecting harmful effects of toxins on the functionality and morphology of the organoids [161, 162], and the ability of organoids to be derived from tumorigenic tissue for drug testing [163]. Nevertheless, these studies have not yet been investigated in iPSC-derived organoids.

CRISPR/Cas9 (clustered regularly interspaced short palindromic repeats-associated protein 9) mediated gene editing has been applied to iPSC research to correct or induce genetic mutations in iPSC lines, primarily in the
study of monogenic diseases. Figure 3 demonstrates the potential research and clinical applications of CRISPR/Cas9-edited iPSCs in domestic species. Extensive research has been done in human models and has been reviewed previously [164, 165]. Although the use of genome editing technologies has been limited in domestic species, a single report of successful CRISPR editing of bovine iPSCs has been published [166]. Genome editing of iPSCs in combination with chimera generation provide the potential for transgenic animal development. In agricultural animals, the artificial introduction of valuable traits e.g. therapeutic proteins in milk, decreased waste product, and disease resistance, could be invaluable to the farming industry [20, 167]. Economically, this would require germline transmission, which has seen little success in domestic species as compared to rodents [20, 116, 168]. Further research is required to fully understand the feasibility, safety and ethical implications of germline transmission of genetically modified iPSCs from domestic species.

Proteomic, metabolic, and methylomic analysis of iPSCs have limited acknowledgement and investigation in research. However, there have been recent efforts to investigate the proteins and sites of methylation of human iPSCs and their derivatives. The investigation of -omics in iPSCs can assist in confirming how completely iPSCs have been reprogrammed to an undifferentiated stem cell state and resemble ESCs. Studying proteins and sites of methylation have clinical applications in autologous cell replacement therapy [169]. Aside from one study investigating the effects of epigenetic modifiers on silencing on exogenous transcription in piPSCs [65], epigenetics is an unexplored area of iPSC research in domestic species. In humans, research has shown isogenic iPSC populations and similar epigenetic markers of hiPSCs and hESCs [170]. Such results further emphasize the potential applicability of iPSCs in disease research and as a substitute for ESCs.

**Fig. 3** Potential Use of Domestic Animal iPSCs for Drug Discovery, Disease Modelling and Cell Replacement Therapy. Induced pluripotent stem cell (iPSCs) can be generated from healthy animals (e.g. dogs) for allogeneic cell transplantation of therapeutic cell types/tissue indicative of the disease. Alternatively, iPSCs can be generated from animals harbouring a genetic disorder and through CRISPR/Cas9-mediated genome editing technologies these genetic mutations can be corrected so that differentiated cell products from these iPSCs can be utilized in autologous cell replacement therapies. In addition, both the genetically mutated iPSCs and their genome-corrected iPSCs can be compared and contrasted for disease modelling purposes. This disease-in-a-dish could be potentially use as a high throughput screening system to discover novel drug candidates. Figure by Dean H. Betts (Adobe Photoshop)
Although there is a steadily growing number of publications pertaining to porcine, equine, and canine models, the numbers are much fewer for cattle, goats, chickens, and cats. Hence, new research initiatives should further investigate these species for their potential application in the fields of disease modelling, treatment, and enhancement of production animals.

Conclusion
Induced pluripotent stem cells are an innovative tool that hold great potential in contributing to veterinary medicine. Protocols for the production of iPSCs in some domestic species have been well-defined and have prompted research into their many potential applications. iPSC cultures have allowed for the production of tissues that can be studied for their physiological use and disease pathologies. Further, iPSCs themselves may be used in the future for the treatment of various diseases seen by veterinary practitioners. Although achievements have been made, a great deal of work is still required before these techniques can be clinically applied.

Abbreviations
- iPSCs: Induced pluripotent stem cells
- piPSCs: Porcine induced pluripotent stem cells
- VSMCs: Vascular smooth muscle cells
- NPCs: Neural progenitor cells
- MVEV: Murray Valley encephalitis virus
- iPSCs: Induced pluripotent stem cells
- MSCs: Mesenchymal stromal cells
- ESCs: Embryonic stem cells
- OSKM: Oct4, Sox2, Klf4, and c-Myc
- ciPSCs-ECs: Canine induced pluripotent stem cells – endothelial cells
- ePSCs: Equine induced pluripotent stem cells
- ciPSCs: Canine induced pluripotent stem cells
- eiPSCs: Equine induced pluripotent stem cells

Acknowledgements
Not applicable.

Authors’ contributions
RS and SP contributed to the conception of the review, data collection and analysis, and drafting of the manuscript. TK and KR contributed to the conception of the review. RS, SP, TK, KR, and DBs contributed to the manuscript revision. RS, SP, TK, KR, and DB read and approved the final manuscript.

Author’s information
Not applicable.

Funding
Dr. Koch’s work is supported by numerous granting agencies including Equine Guelph, Per-Trust, Natural Sciences and Engineering Research Council of Canada, Morris Animal Foundation, Canadian Foundation of Innovation, and the Ontario Ministry of Innovation.

Availability of data and materials
All data analyzed during this study is included in this published article.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1Department of Biomedical Sciences, Ontario Veterinary College, University of Guelph, 50 Stone Road East, Guelph, Ontario N1G 2W1, Canada. 2Department of Physiology and Pharmacology, The University of Western Ontario, London, Ontario N6A 5C1, Canada.

Received: 2 July 2020 Accepted: 24 November 2020
Published online: 08 December 2020

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