

ENCYCLOPEDIA

of

CHEMICAL

TECHNOLOGY

Fourth Edition

VINYL POLYMERS (PVC)

POLY(VINYL CHLORIDE)

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September 16, 1996

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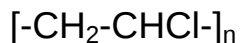
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INTRODUCTION

Poly(vinyl chloride) [9002-86-2] (PVC) has large and broad uses in commerce. It is second in volume only to polyethylene and had a volume sales in North America in 1995 of 6.2 billion kilograms (13.7 billion pounds) (1). See table 1 for volumes of several thermoplastics. This large volume of sales can be attributed to several unique properties. Vinyl compounds usually contain close to 50% chlorine. Not only does the chlorine provide no fuel, it acts to inhibit combustion in the gas phase, providing the vinyl with a very high level of combustion resistance, useful in many building applications as well as many electrical housings and electrical insulation applications.

PVC has a unique ability to be compounded with a wide variety of additives, making it possible to produce materials in a range from flexible elastomers to rigid compounds, materials that are virtually unbreakable with a notched Izod impact greater than 0.5 J/mm at -40°C, materials that are weatherable with good property retention for over 30 years, compounds that have stiff melts and little elastic recovery for outstanding dimensional control in profile extrusion, or low viscosity melts for thin walled injection molding.

Produced by free radical polymerization, PVC has the following structure:



where the degree of polymerization, n , ranges from 500 to 3500.

The first discovery of PVC was in 1872 when E. Baumann found that exposure of vinyl chloride to sunlight produced a white solid that resisted attack by potassium hydroxide or water and melted with degradation at above 130°C. (2, 3). From 1912 to 1926, German workers at Chemische Fabrik Griesheim-Electron tried but failed to build machinery that could process PVC and overcome its instability; finally they gave up their patents (3). In 1926, Waldo Semon at BFGoodrich, while looking for an adhesive to bond rubber to metal for tank liners, found that boiling PVC in tricresyl phosphate or dibutyl phthalate, made it highly elastic (3), thus he invented the first thermoplastic elastomer.

Table 1. 1995 SALES VOLUME IN NORTH AMERICA (1).
 BILLIONS OF KILOGRAMS

	USA	CANADA	MEXICO	NORTH AMERICAN TOTAL
PVC, POLY(VINYL CHLORIDE)	5.36	0.45	0.40	6.22
HDPE, HIGH DENSITY POLYETHYLENE	5.29	0.73	0.18	6.20
LLDPE, LINEAR LOW DENSITY POLYETHYLENE	2.70			2.70
LDPE, LOW DENSITY POLYETHYLENE	3.38	1.31	0.30	4.99
PP, POLYPROPYLENE	4.84	0.26	0.22	5.32
PS, POLYSTYRENE	2.68		0.14	2.81
ABS, ACRYLONITRILE/ BUTADIENE/STYRENE	0.66			0.66
THERMOPLASTIC POLYESTER	1.78			1.78
PC, POLYCARBONATE	0.34			0.34
ACETAL	0.15			0.15
ACRYLIC	0.25			0.25
NYLON	0.47			0.47

DISCUSSION

PHYSICAL PROPERTIES

Morphology as polymerized

The major type of polymerization of PVC is the suspension polymerization route. The morphology formed during polymerization strongly influences the processability and physical properties. Mass polymerized PVC has a similar morphology to suspension PVC.

In the suspension polymerization of PVC, droplets of monomer 30 - 150 μm in diameter are dispersed in water by agitation. A thin membrane is formed at the water/monomer interface by dispersants such as polyvinyl alcohol or methyl cellulose. This membrane, isolated by dissolving the PVC in tetrahydrofuran and measured at 0.01 - 0.02 μm thick, has been found to be a graft copolymer of polyvinyl chloride and polyvinyl alcohol (4, 5). Early in the polymerization, particles of PVC deposit onto the membrane from both the monomer and the water sides forming a skin 0.5 - 5 μm thick that can be observed on grains sectioned after polymerization (4, 6). Primary particles, 1 μm diameter, deposit onto the membrane from the monomer side and water phase polymer, 0.1 μm diameter, deposits onto the skin from the water side of the membrane (4). These

domain sized water phase particles may be one source of the observed domain structure (7).

Figure 1. A grain of suspension PVC and its cross-section showing the skin and primary particles.

Grain
~ 150 μm dia.

Skin
~ 2-5 μm thick

Primary particles
~ 1 μm dia.

Figure 2. Skin of a suspension PVC grain showing 0.1 mm diameter particles deposited from the water phase.

Mass polymerized PVC also has a skin of compacted PVC primary particles very similar in thickness and appearance to the suspension polymerized PVC skin, compared in figure 3, however, mass PVC would not contain the thin block copolymer membrane (7).

Figure 3. Cross-sections of PVC grains from a) suspension and b) mass PVC grains showing similar morphologies.

mass

suspension

In suspension PVC polymerization, droplets of polymerizing PVC, 30 - 150 μm diameter, agglomerate to form grains at 100 - 200 μm diameter (8). With one droplet per grain, the shape is quite spherical. With several droplets making up the grain, the shape can be quite irregular and knobby (9). The grain shape plays an important role in determining grain packing and bulk density of a powder (9).

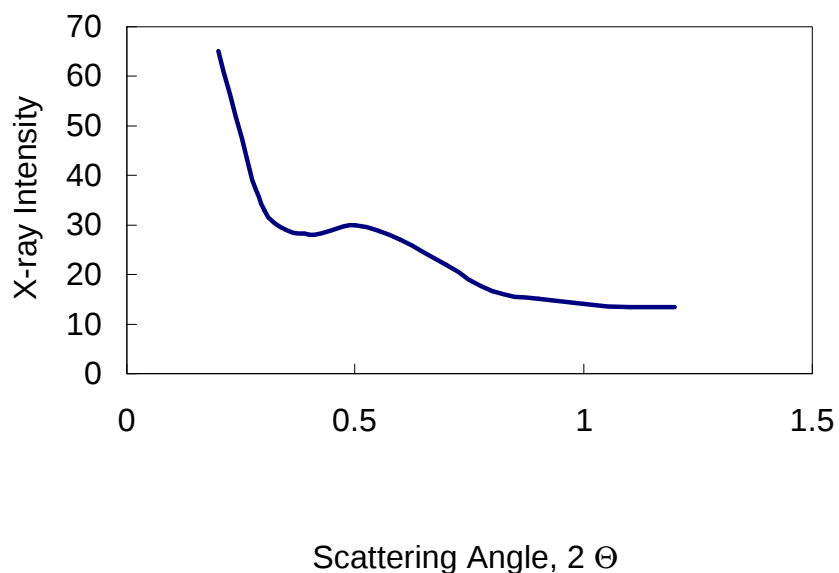
For both suspension and mass polymerizations at less than two percent conversion, PVC precipitates from its monomer as stable primary particles, slightly below 1 μm diameter (4, 10 - 12). These primary particles are stabilized by a negative chloride charge (4, 13). Above two percent conversion, these

primary particles agglomerate. Sectioning the PVC grains of either suspension or mass resins readily shows the skins, primary particles at 1 μm diameter, and agglomerates of primary particles at 3 - 10 μm diameter (4, 7, 8, 14).

These primary particles also contain smaller internal structures. Electron microscopy reveals a domain structure at about 0.1 μm diameter (8, 15, 16). The origin and consequences of this structure is not well understood. As mentioned earlier, PVC polymerized in the water phase and deposited on the skin, may be the source of some of the domain sized structures. Also domain sized flow units may be generated by certain unusual and severe processing conditions, such as high temperature melting at 205°C followed by lower temperature mechanical work at 140 - 150°C.(17), that break down the primary particles further.

On an even smaller scale is the microdomain structure at 0.01 μm spacing. Small angle x-ray scattering reveals a scattering peak (figure 4) corresponding to density fluctuations spaced at about 0.01 μm (18 - 20). When PVC is swelled by plasticizer or swelled with a poor solvent such as acetone, swelling reaches a limit where the PVC will no longer absorb more plasticizer or acetone (21). This data suggests a structure where the crystallites of about 0.01 μm spacing are tied together by molecules in the amorphous regions. Plasticizer or acetone only swells the amorphous regions without dissolving the crystallites. Also electron microscopy shows a spacing in plasticized PVC of 0.01 μm (22).

Figure 4. Small angle X-ray scattering pattern from PVC plasticized with 20 parts per hundred resin of dioctyl phthalate (18).



Hierarchical structure of PVC

As we have just discussed, PVC has structure which is built upon structure which is built upon even more structure. These many layers of structure are all important to performance and are inter-related. A summary of these structures is listed in table 2. Also figure 5 examines a model of these hierarchies on three scales.

Table 2. Summary of polyvinyl chloride morphology.

<i>Feature</i>	<i>Size</i>	<i>Description</i>
Droplets	30 - 150 μm diameter	The dispersed monomer during suspension polymerization.
Membranes	0.01 - 0.02 μm thick	The membrane at the monomer-water interface in suspension PVC. It is usually a graft copolymer of PVC and the dispersant such as polyvinyl alcohol.
Grains	100 - 200 μm diameter	After polymerization, the free flowing powder that is usually made up of agglomerated droplets. In mass polymerization it is the free flowing powder.
Skins	0.5 - 5 μm thick	The shell on grains made up of PVC deposited onto the membrane during suspension polymerization. In mass polymerization it is PVC compacted on the surface of the grain.
Primary particles	1 μm diameter	Formed as a single polymerization site in both suspension and mass polymerization by precipitation of polymer from the monomer. Made up of a billion molecules, it is often the melt flow unit established during melt processing. In emulsion polymerization it is the emulsion particle.
Agglomerates of primary particles	3 - 10 μm diameter	Formed during polymerization by the merging of primary particles.
Domains	0.1 μm diameter	Formed under special conditions such as high temperature melting at 205°C followed by lower temperature mechanical work at 140 - 150°C. Water phase polymerization also produces domain sized structure.
Microdomains	0.01 μm spacing	Crystallite spacing.
Secondary crystallinity	0.01 μm spacing	Crystallinity that is re-formed from the amorphous melt and is responsible for fusion (gelation).

Figure 5. The hierarchical structure of PVC.

Morphology during processing

The first step in processing is usually powder mixing in a high speed, intensive mixer. PVC resin, stabilizers, plasticizers, lubricants, processing aids, fillers, and pigments are added to the powder blend for distributive mixing. For both suspension and mass PVC resins, intensive mixing above the glass transition temperature, results in a progressive increase in apparent bulk density with higher mixing temperatures (23). This increase in apparent bulk density is due to the smoothing and rounding of the irregular surface. However the grains of PVC are largely unchanged and are not grossly deformed nor are they broken down to smaller particles (23).

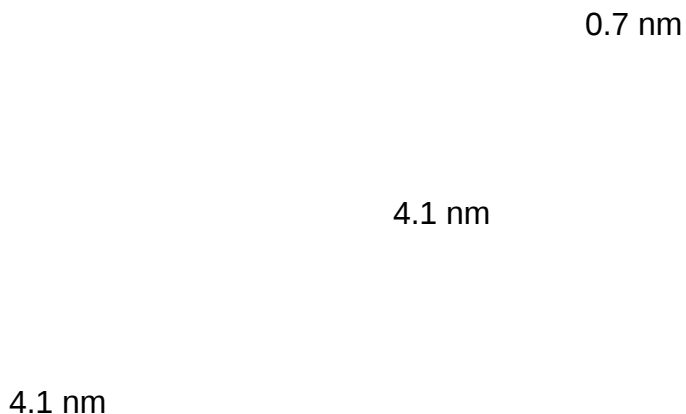
In plasticized PVC, liquid plasticizers first fill the voids or pores in the PVC grains fairly rapidly during powder mixing. If a large amount of plasticizer is added, the excess plasticizer beyond the capacity of the pores, initially remains on the surface of the grains making the powder somewhat wet and sticky. Continued heating increases the diffusion rate of plasticizer into the PVC mass where the excess liquid is eventually absorbed and the powder dries.

PVC powder compounds are heated, sheared, and deformed during melt processing. During this process, the grains of PVC are broken down. First the skin is torn, exposing the PVC grain's internal structures (24). Then the grains are broken down to agglomerates of primary particles, then to primary particles

as the melt flow units. The primary particles seem to be persistent and fairly stable structures in the melt (25 - 32). This processing window of stable primary particles exists even with continued melt processing. The primary particle is about a billion molecules of PVC held together by a structure of crystallites and tie molecules (21).

The PVC crystallites are small (average 0.7 nm = 3 monomer units) in the PVC chain direction, and are packed laterally to a somewhat greater extent (4.1 nm) (21, 33). A model of the crystallite is drawn in figure 6. The crystalline structure of PVC is found to be an orthorhombic system, made of syndiotactic structures, two monomer units per unit cell, and 1.44 - 1.53 specific gravity (34 - 37).

Figure 6. The crystallite structure of PVC.



PVC fusion (gelation)

The PVC primary particle flow units (billion molecule bundles) can partially melt, free some molecules of PVC, which can entangle at the flow unit boundary. These entangled molecules can recrystallize upon cooling, forming secondary crystallites, and tie the flow units together into a large three dimensional structure (21, 38). This process is known as fusion or gelation.

The strength created by the fusion process is strongly dependent on the previous processing temperature and the molecular weight of the PVC (38 - 44). The degree of fusion or gelation is measured in several ways. The entrance pressure in capillary rheometry is often used as a measure of the fusion strength

(40, 42, 45 - 53). Differential scanning calorimetry also is often used as an indication of previous melt temperature and the amount of crystallinity melted and reformed as secondary crystallinity (54, 55). X-ray defraction has been used to measure gelation (55). Acetone or methylene chloride swelling and observation of structural break-down is widely used as a qualitative measure of fusion (44, 56 - 59). Sometimes the acetone swollen specimens are sheared between glass slides to further establish the strength of the three dimensional structure (59). Fusion has also been assessed based on scanning electron microscopy of fractured surfaces (29, 30, 46). And inverse gas chromatographic measurements have also been useful in accessing the degree of fusion of PVC (60).

The strength of this large three dimensional fused (gelled) structure has been shown to be critical in determining Izod impact, creep rupture strength, and even flow in rigid injection molding. In these cases both the melt temperature during processing and the PVC molecular weight play a large role in the Izod and creep rupture (38 - 41). In plasticized PVC, this large three dimensional structure (also dependent on molecular weight and previous processing temperature) determines tensile strength, creep, and cut resistance. A model for accounting for molecular weight effects and processing temperature effects on PVC fusion is presented in figure 7.

Figure 7. Model for PVC fusion, accounting for molecular weight effects and processing temperature effects.

a) Unfused PVC primary particles.

b) Partially melted PVC primary particles.

c) Partially melted then recrystallized high molecular weight PVC, showing strong three dimensional structure.

d) Partially melted then recrystallized low molecular weight PVC, showing weak three dimensional structure.

PVC normally improves in properties with increasing fusion (or increasing melt temperature), however some observations show a fall off in impact properties with higher melt temperatures (42, 56). This has been shown to be caused by melt fracture, when PVC fuses and flows as large melt flow units of multiple primary particles (61). These flow units of fused multiple primary particles can lead to surface roughness during extrusion, in both rigid and plasticized PVC (44, 56, 62); they can also be responsible for roughness when improperly handling regrind (63).

Plasticized PVC Morphology

While most of the discussion has been on rigid PVC, plasticized PVC has the same structures as rigid PVC, except that plasticizer enters the amorphous phase of PVC making the tie molecules elastomeric. The grains break down to 1 μm primary particles which become the melt flow units (44). The crystallites are not destroyed by plasticizer (21). Partial melting allows entanglement at the flow unit boundaries, followed by recrystallization upon cooling to form a strong three dimensional elastomeric structure (38, 45).

PVC physical parameters

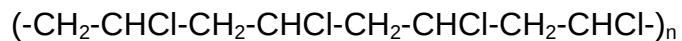
Table 3. PVC physical parameters.

PVC PROPERTY	VALUE	REFERENCE
Crystallographic data	orthorhombic 2 monomer units/cell	
	<u>a</u> <u>b</u> <u>c</u>	
commercial PVC	1.06 0.54 0.51 nm	34
single crystal	1.024 0.524 0.508 nm	37
Percent crystallinity		
as polymerized	19%	
from the melt	4.9%	64
Density (uncompounded)		
whole	1.39 grams/cc.	65
crystallites	1.53 grams/cc.	37
Oxygen permeability		
	$238 e^{-13.3/RT}$	66
	cc/cm sec cm ² cm Hg	
Poisson ratio (rigid PVC)	0.41	
Refractive index	1.54	67
Glass transition temperature	83°C.	
Coefficient of linear thermal expansion (unplasticized)	$7 \times 10^{-5}/^{\circ}\text{C}.$	
Specific heat	<u>temp.</u> <u>value</u>	
rigid PVC	23°C. 0.22 cal./gram °C.	68
	50 0.25	
	80 0.35	
	120 0.39	
plasticized PVC (50 phr DOP)	23 0.37	68
	50 0.40	
	80 0.42	
	120 0.45	
Thermal conductivity (unplasticized)	4.2×10^{-4} cal/cm sec °C.	69
Dielectric strength	0.5 Kvolts/mil	
	20 Kvolts/mm.	
Solubility parameter	9.75 (average)	70
	[cal./cm ³] ^{0.5}	

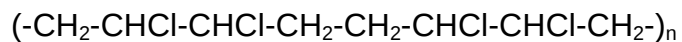
CHEMICAL PROPERTIES

Molecular structure - Monomer addition orientation

The addition of vinyl monomer to a growing PVC chain can be considered to add in a head-to-tail fashion, resulting in a chlorine atom on every other carbon atom,



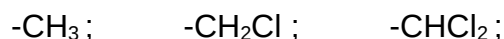
or in a head-to-head, tail to tail fashion, resulting in chlorine atoms on adjacent carbon atoms.



Dechlorination of head-to-head, tail-to-tail structure would be expected to go to 100% completion. If dechlorination of head-to-tail structure starts at random positions, then 13.5% of the chlorine should remain at the end of reaction. Dilute solutions of PVC treated with zinc, removes 87% of the chlorine, proving the head-to-tail structure of PVC (71).

End groups and branching

Both saturated and unsaturated end groups can be formed during polymerization by chain transfer to monomer or polymer and by disproportionation. Some of the possible chain end groups are shown:

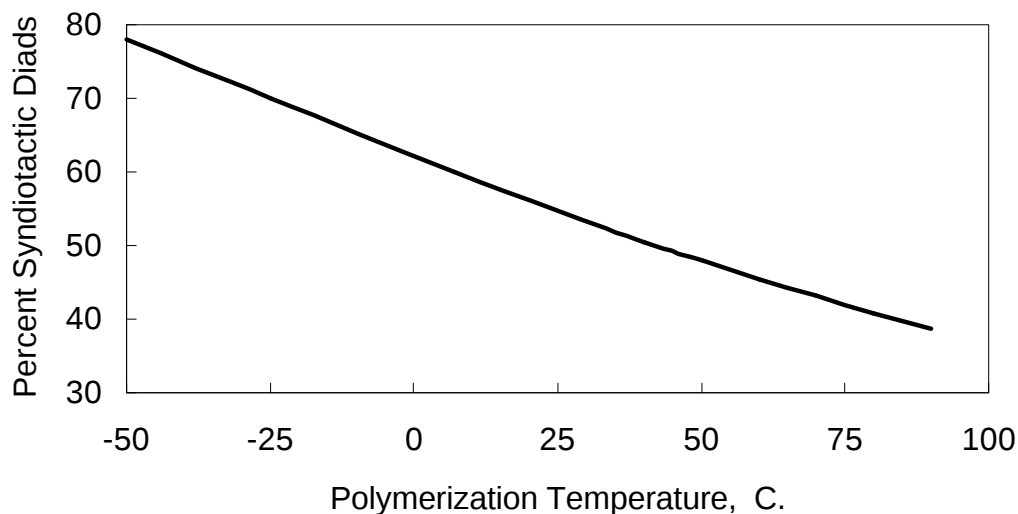


PVC polymerization has a high chain transfer activity to monomer; about 60% of the chains have unsaturated chain ends (72) and the percentage of chain ends containing initiator fragments is low (73). Chain transfer to polymer leads to branching. Branching in PVC has been measured by hydrogenating PVC, removing chlorine with lithium aluminum hydride. The ratio of methyl to methylene groups is measured by infrared spectroscopy using bands at 1378 and 1350, or 1370 and 1386 cm^{-1} . Conventional PVC resins, made by mass or suspension polymerization at 50 - 90°C, contain 0.2 to 2 branches per 100 carbon atoms (65, 73).

Stereoregularity

The addition of monomer fixes the tacticity of the previous monomer unit. Syndiotactic structure has the adjacent chlorine atoms oriented to opposite sides of the carbon-carbon-carbon plane. Isotactic structure has the adjacent chlorine atoms oriented to same side of the carbon-carbon-carbon plane. The potential energy for syndiotactic conformation is 4.2 - 8.4 kJ/mol (1-2 kcal/mol) lower than for isotactic conformation (74, 75). Then the ratio of propagation rates for syndiotactic to isotactic, k_s/k_i , must increase with decreasing temperature. Thus with decreasing polymerization temperature, the degree of syndiotacticity in PVC should increase. Measured amounts of syndiotacticity are illustrated in figure 8 (76, 77).

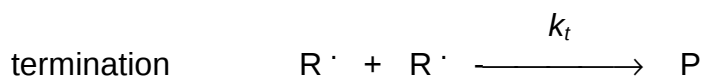
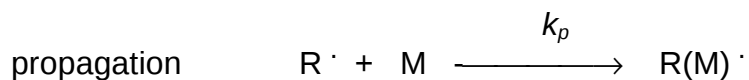
Figure 8. The Syndiotactic Structure of PVC.



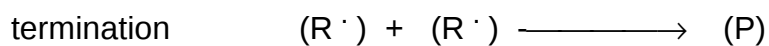
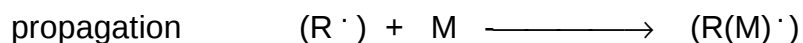
Polymerization kinetics of mass and suspension PVC

The polymerization kinetics of mass and suspension PVC are considered together because a droplet of monomer in suspension polymerization can be considered to be a mass polymerization in a very tiny reactor. During polymerization, the polymer precipitates from the monomer when the chain size reaches 10 - 20 monomer units. The precipitated polymer remains swollen with monomer, but has a reduced radical termination rate. This leads to a higher concentration of radicals in the polymer gel and to an increased polymerization rate at higher polymerization conversion.

Reactions in the liquid phase:



Reactions in the polymer gel:

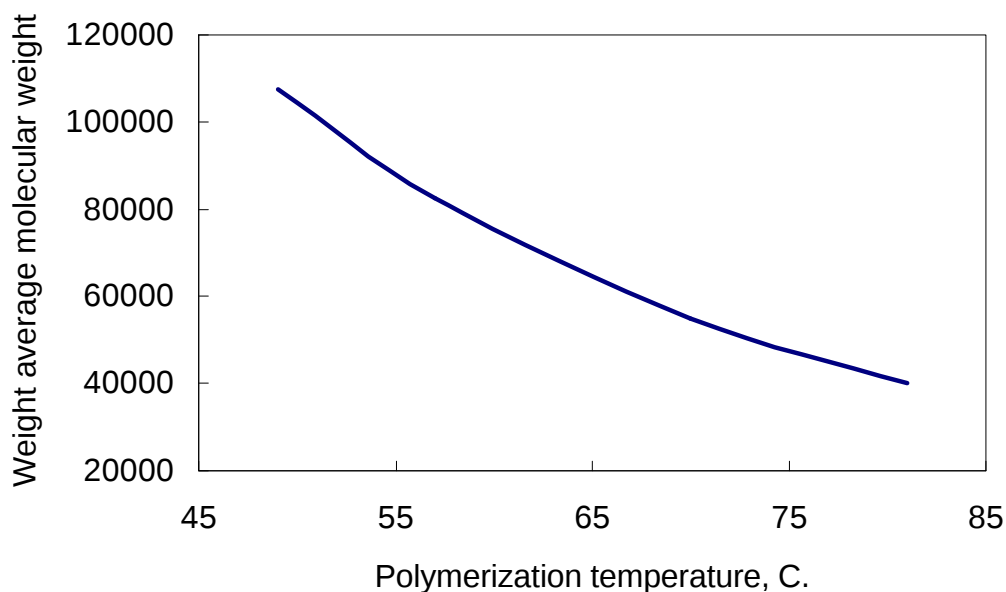


where $R^\cdot$ = a polymer chain radical in liquid monomer, $(R^\cdot)$ = a polymer chain radical in the polymer gel phase, M = monomer molecule, (M) = monomer molecule in the polymer gel phase, P = polymer in monomer, (P) = polymer gel, and k_i , k_p , and k_t are reaction rate coefficients for initiation, propagation and termination. Values for k_p , and k_t at 60°C are 1.23×10^5 and 2.3×10^{10} L/(mol sec), respectively (78).

Polymerization in two phases, the liquid monomer phase and the swollen polymer gel phase, form the basis for kinetic descriptions of PVC polymerization (79 - 81). The polymerization rate is slower in the liquid monomer phase than in the swollen polymer gel phase due to the greater mobility in liquid monomer which allows for greater termination efficiency. The lack of mobility in the polymer gel phase, reduces termination, creating a higher concentration of radicals, and thus creating a higher polymerization rate. Thus the polymerization rate increases with conversion to polymer.

Chain transfer to monomer is the main reaction controlling molecular weight and molecular weight distribution. The chain transfer constant to monomer, C_m , is the ratio of the rate coefficient for transfer to monomer to that of chain propagation. $C_m = 6.25 \times 10^{-4}$ at 30°C and 2.38×10^{-3} at 70°C and a general expression is $C_m = 5.78 e^{-2768/T}$. At 30°C, chain transfer to monomer happens once in every 1600 monomer propagation reactions and at 70°C, chain transfer happens once every 420 monomer additions (80, 82 - 84). Thus temperature of polymerization strongly influences PVC molecular weight, the molecular weight increasing with lower polymerization temperature as indicated in figure 9.

Figure 9. PVC molecular weight as influenced by polymerization temperature.



PVC molecular weights are usually determined in the USA using inherent viscosity or relative viscosity measured according to ASTM D1243, 0.2 grams/100 ml of cyclohexanone at 30°C. In Europe, K values are used, measured at 0.5% in cyclohexanone. The relationship between inherent viscosity, K value, number average molecular weight (M_n), and weight average molecular weight (M_w) for commercial grades of PVC are shown in table 4 (85).

Table 4. Quick reference for commercial PVC molecular weights.

Inherent viscosity, ASTM D1234	Relative viscosity, ASTM D1234	K value, (Fikentscher: DIN 53726)	Number average molecular weight M_n ($\times 10^{-3}$)	Weight average molecular weight M_w ($\times 10^{-3}$)
0.42	1.09	45.0	15.0	30.0
0.47	1.10	47.1	18.0	36.0
0.52	1.11	49.3	20.0	40.0
0.57	1.12	51.3	22.5	45.0
0.62	1.13	53.6	25.0	50.0
0.67	1.14	56.1	27.5	55.0
0.73	1.16	58.2	30.5	61.0
0.78	1.17	60.5	33.0	67.0
0.83	1.18	62.9	36.0	72.0
0.88	1.19	64.9	38.5	78.0
0.92	1.20	67.1	41.0	82.5
0.98	1.22	69.2	44.0	89.5
1.03	1.23	71.5	47.0	95.0
1.08	1.24	73.3	50.0	101.0
1.13	1.25	74.9	52.5	107.5
1.21	1.27	77.5	57.0	117.0
1.30	1.30	80.7	62.5	128.5
1.40	1.32	83.8	68.5	141.0
1.60	1.38	90.8	81.0	168.0
1.80	1.43	96.7	93.5	195.0

PVC RESIN MANUFACTURING PROCESSES

Mass Polymerization

Mass or bulk polymerization of PVC is normally difficult. At high conversions the mixture becomes extremely viscous, impeding agitation and heat removal, causing a high polymerization temperature and broad molecular weight distribution (86). A two stage process that overcomes these problems was originally developed by Saint Gobain (France). The first stage of the process is carried out in a prepolymerizer with flat blade agitator and baffles to about 7 - 10% conversion. This first stage forms a skeleton seed grain for polymerization in a second stage. The number of grains remain constant throughout this polymerization (87).

In the second stage polymerizer, a larger horizontal vessel, more monomer and initiator are added. This vessel is equipped with a slow moving agitator blade running close to the vessel wall. The reaction proceeds through the liquid stage and at about 25% conversion, becomes a powder. Heat removal is achieved, 30% by the jacket, 60% by a condenser, and 10% by the cooled agitator shaft (88). Unreacted monomer is removed by vacuum. While the mass process saves drying energy, it has remained a minor process when compared to the suspension process.

Suspension Polymerization

Suspension polymerization is carried out in small droplets of monomer suspended in water. The monomer is first finely dispersed in water by vigorous agitation. Suspension stabilizers act to minimize coalescence of droplets by forming a coating at the monomer-water interface. The hydrophobic-hydrophilic properties of the suspension stabilizers are key to resin properties and grain agglomeration (89).

Kinetics of suspension PVC are identical to the kinetics of mass PVC, increasing in rate with conversion (90). After polymerization to about 80-90% conversion, excess monomer is recovered, the slurry is steam stripped in a column to a residual monomer level of about 0.0001 % (10 ppm), excess water is centrifuged off, and the resin is dried with hot air.

Emulsion Polymerization

Emulsion polymerization takes place in a soap micelle where a small amount of monomer dissolves in the micelle. The initiator is water soluble. Polymerization takes place when the radical enters the monomer swollen micelle (91, 92). Additional monomer is supplied by diffusion through the water phase. Termination takes place in the growing micelle by the usual radical-radical interactions. A theory for true emulsion polymerization postulates that the rate is proportional to the number of particles $[N]$; N depends on the 0.6 power of the

soap concentration $[S]$ and the 0.4 power of initiator concentration $[I]$; the average number of radicals per particle is 0.5 (93).

However, the kinetics of PVC emulsion does not follow the above theory. The rate shows the same increasing behavior with conversion as mass polymerization (94, 95). $[N]$ depends on $[S]$, but the relationship varies with the emulsifier type (96, 97). However the rate is nearly independent of $[N]$ (95). The average number of radicals per particle is low, 0.0005 to 0.1 (95). The high solubility of vinyl chloride in water, 0.6 weight %, accounts for a strong deviation from true emulsion behavior. Also PVC's insolubility in its own monomer accounts for some behavior such as a rate dependence on conversion.

Emulsions of up to 0.2 μm diameter are sold in liquid form for water based paints, printing inks, and finishes for paper and fabric. Other versions, 0.3 to 10 μm diameter and dried by spray drying or coagulation, are used as plastisol resins. Plastisols are dispersions of PVC in plasticizer. Heat allows fast diffusion of plasticizer into the PVC particle followed by fusion (gelation) to produce a physically crosslinked elastomer, where the physical crosslinks are PVC crystallites.

Micro-Suspension Polymerization

While emulsion polymerization uses a water soluble initiator, micro-suspension polymerization uses a monomer soluble initiator. The monomer is homogenized in water along with emulsifiers or suspending agents to control the particle sizes. Micro-suspension paste resins at 0.3 to 1 μm diameter are used to make plastisols for flooring, seals, barriers, etc. These plastisols are also dispersions of PVC in liquid plasticizer and are cured by heating. Heating allows plasticizer to uniformly diffuse into the PVC particles and at higher temperatures, the plasticized particles fuse. Micro-suspension blending resins at 10 to 100 μm diameter are used as extenders to paste resins in plastisols (98).

Solution Polymerization

In solution polymerization a solvent for the monomer is often used to obtain very uniform copolymers. Polymerization rates are normally slower than for suspension or emulsion PVC. For example, vinyl chloride, vinyl acetate, and sometimes maleic acid are polymerized in a solvent where the resulting polymer is insoluble in the solvent. This makes a uniform copolymer, free of suspending agents, that is used in solution coatings (99).

Copolymerization

Vinyl chloride can be copolymerized with a variety of monomers. Vinyl acetate is the most important commercial comonomer [9003-22-9]. It is used to reduce crystallinity which aids fusion and allows lower processing temperatures. Copolymers are used in flooring and coatings. This copolymer sometimes contains maleic acid or vinyl alcohol (hydrolyzed from the polyvinyl acetate) to improve the coating's adhesion to other materials including metals. Copolymers with vinylidene chloride are used as barrier films and coatings. Copolymers of vinyl chloride with maleates or fumarates are used to raise heat deflection temperature. Copolymers of vinyl chloride with acrylic esters in latex form are used as film formers in paint, non-woven fabric binders, adhesives, and coatings. Copolymers with olefins improve thermal stability and improve melt flow at some loss of heat deflection temperature (100). Copolymerization parameters are listed in table 5.

Table 5. Copolymerization Parameters of Vinyl Chloride.

M ₂	r ₁	r ₂	e	Q	temperature °C.	ref.
acrylic acid	0.107	6.8	0.77	1.15	60	101
acrylonitrile	0.04	2.7	1.20	0.60	60	102
butadiene	0.035	8.8	-1.05	2.39	50	103
butene	3.4	0.21				104
<i>n</i> -butyl acrylate	0.07	4.4	1.06	0.50	45	101
						105
diethyl fumarate	0.12	0.47	1.25	0.61	60	106
dimethyl itaconate	0.053	5.0	1.34	1.03	50	107
diethyl maleate	0.77	0.009				104
ethylene	3.21	0.21	-0.20	0.015	50	108
ethylhexyl acrylate	0.16	4.15				104
isobutylene	2.05	0.08	-0.96	0.033	60	109,
						110
isoprene			-1.22	3.33		
maleic anhydride	0.296	0.008	2.25	0.23	75	111
methacrylic acid	0.034	23.8	0.65	2.34	60	101
methacrylonitrile			0.68	0.86	60	112
methyl acrylate	0.12	4.4	0.60	0.42	50	113
methyl methacrylate	0.1	10	0.4	0.74	68	114
octyl acrylate	0.12	4.8	1.07	0.35	45	105
propylene	2.27	0.3	-0.78	0.002		115,
						110
styrene	0.02	17	-0.80	1.0	60	116
vinyl acetate	1.68	0.23	-0.22	0.026	60	117
N-vinylcarbazole	0.17	4.8	-1.40	0.41	50	118
vinyl chloride			0.20	0.044		
vinyl laurate	7.4	0.2				104
vinylidene chloride	0.3	3.3	0.36	0.22	60	119
vinyl isobutyl ether	2.0	0.02			50	120
N-vinylpyrrolidone	0.53	0.38	-1.14	0.14	50	121

COMPOUNDING

The additives found in PVC help make it one of the most versatile, cost-efficient materials in the world. Without additives, literally hundreds of commonly used PVC products would not exist. Many materials are useless until they undergo a similar modification process. Steel, for instance, contains among other things, chromium, nickel and molybdenum. PVC's are tailored to the requirements using sophisticated additives technology.

Stabilizers

Lead stabilizers, particularly tribasic lead sulfate is commonly used in plasticized wire & cable compounds because of its good non-conducting electrical properties (122).

Organotin stabilizers are commonly used for rigid PVC, including for pipe, fittings, windows, siding profiles, packaging and injection molded parts.. These repair unstable sites on PVC removing unstable chlorine and replacing it with a ligand from the tin stabilizer molecule(123 - 125). This produces stability at least an order of magnitude better than without stabilizer. Examples of effective tin compounds are di-alkyl tin dilaurate, and mono-/di-alkyl tin di-isooctylthioglycolate. Certain grades of methyl tins and octyl tins are used in food contact applications.

Antimony tris(isooctylthioglycolate) has found use in pipe formulations at low levels. Its disadvantage is it cross-stains with sulfide based tin stabilizers(122).

Barium-zinc stabilizers have found use in plasticized compounds, replacing barium-cadmium stabilizers. These are used in moldings, profiles, and wire coatings. Cadmium use has decreased because of environmental concerns surrounding certain heavy metals.

Calcium-zinc stabilizers are used in both plasticized PVC and rigid PVC for food contact where it is desired to minimize taste and odor characteristics. Applications include meat wrap, water bottles, and medical uses.

Many stabilizers require co-stabilizers. Several organic co-stabilizers are quite useful with barium-zinc and calcium-zinc stabilizers. They are beta diketones, epoxies, organophosphites, hindered phenols, and polyols (122).

Impact modifiers

In the early days of plastics, many unplasticized PVC products were brittle. This gave plastics a cheap reputation. It was therefore quite desirable to develop technology to produce tough plastics. In the early 1950s, The Geon Company (then a part of BFGoodrich) began adding rubbery polymers to PVC to improve toughness (126). Rubbery particles act as stress concentrators or multiple weak points, leading to crazing or shear-banding under impact load (127). This can result in cavitation and/or cold drawing, thus allowing the PVC to absorb large amounts of energy. Impact modifier choices are listed in table 6.

Table 6. Impact modifiers used for various PVC applications (128).

[illegible]

Processing aids

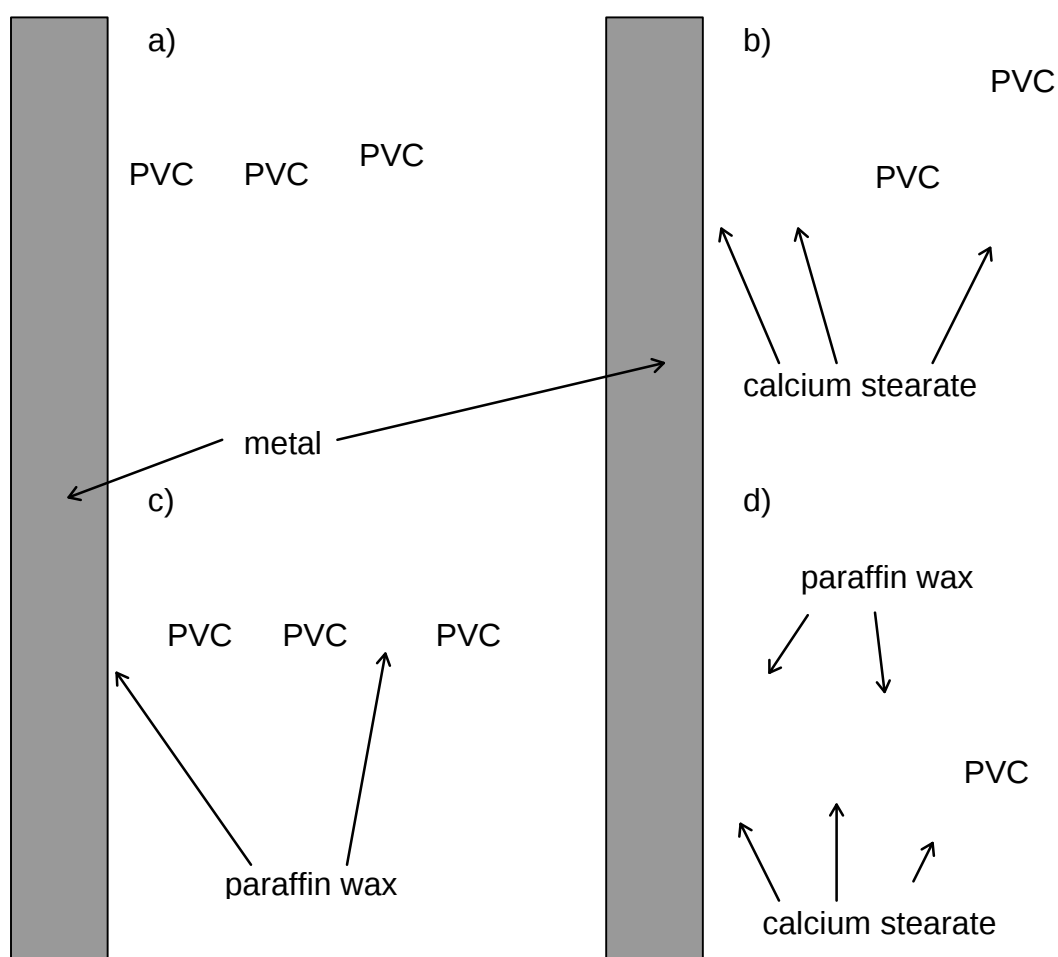
PVC often flows as billion molecule primary particles. Processing aids glue these particles together before the PVC melts, thus acting as a fusion promoter. Processing aids also modify melt rheology by increasing melt elasticity and die swell, or, some reduce melt viscosity, and they reduce melt fracture. Some processing aids lubricate to reduce PVC sticking to metal. And processing aids affect dispersion of fillers, impact modifiers, and pigments (129). The most common processing aids are high molecular weight acrylics based primarily on polymethylmethacrylate copolymers.

Lubricants

Many authors classify lubricants as “internal” or “external” (130 - 134). Internal lubricants were considered to be soluble in PVC, or considered to have little effect on fusion, or they reduce melt viscosity; external lubricants were considered to retard fusion or to promote metal release. This system of classifying lubricants has too many conflicting measurements to be consistent and useful. Others have shown classifications based on synergy between various lubricants (135, 136), but did not explain the nature of that synergy. A model for the lubrication mechanism has been developed which explains synergy between certain lubricants (62). This model treats lubricants as surface active agents. Some lubricants have polar ends which are attracted to other

polar ends and to polar PVC flow units and to polar metal surfaces. These also have non-polar ends which are repelled by the polar groups. Synergy happens when non-polar lubricants are added, which are attracted to the non-polar ends and act as a slip layer. This model is shown in figure 10.

Figure 10. A model of PVC lubrication mechanism (not to scale) showing a) PVC adhesion to metal without lubricant, b) surface activity of calcium stearate, c) non-metal releasing character of just paraffin, and d) synergy between calcium stearate and paraffin (62).



Plasticizers

In 1926 at The Geon Company (then BFGoodrich), solutions of PVC, prepared at elevated temperatures with high boiling solvents, possessed unusual elastic properties when cooled to room temperature (137). Such solutions are flexible, elastic, and exhibit a high degree of chemical inertness and solvent resistance.

This unusual behavior is due to unsolvated crystalline regions in the PVC that act as physical crosslinks, but allow the PVC to accept large amounts of solvent (plasticizers) in the amorphous regions, lowering its T_g to well below room temperature, thus making it rubbery. Thus PVC was the first thermoplastic elastomer (TPE). This rubber-like material has stable properties over a wide temperature range (32, 138 - 140).

A few plasticizers impart specific properties for particular applications. For example, citrate esters are used in food contact applications, benzoates are used for stain resistance, and chlorinated hydrocarbons impart flame resistance and good electrical properties. Aliphatic diesters offer good low temperature flexibility; linear alcohol based phthalates offer good low temperature flexibility and also have reduced volatility; phosphates improve flame resistance; trimellitates have low volatility, are used for high temperature applications, and also have good low temperature properties. Polymeric plasticizers do not migrate easily but suffer from poor low temperature flexibility. Epoxy plasticizers

are also good plasticizers with low volatility and they act as costabilizers,
improving the thermal stability of PVC.

Table 7. Common PVC plasticizers.

Plasticizer	Abbreviation
Aliphatic ester	
di(2-ethylhexyl) adipate	DOA
di(2-ethylhexyl) azelate	
di(2-ethylhexyl) sebacate	
Phthalate	
di(2-ethylhexyl)	DOP or DEHP
diisooctyl	DIOP
diisodecyl	DIDP
butylbenzyl	BBP
butyloctyl	BOP
diisononyl	DINP
ditridecyl	DTDP
diundecyl	DUP
linear C7-C11	711 phthalate
di(2-ethylhexyl) terephthalate	DOTP
Phosphates	
trioctyl	TOP
cresyl diphenyl	CDP
tricresyl	TCP
triphenyl	
tri(2-ethylhexyl)	TEHP
Trimellitates	
tris(2-ethylhexyl)	TOTM
triisooctyl	TIOTM
Epoxies	
epoxidized soybean oil	ESO
epoxidized linseed oil	
epoxy stearate	
2-ethylhexyl epoxytallate	

Plasticizers and stabilizers in particular have been researched at length to determine their potential impact on human health and the environment. DEHP (di-2-ethylhexyl-phthalate) has been used worldwide in applications such as blood bags, saline solutions, meat wraps and other highly credible uses, however, there has been much debate over that impact due to the differing

methods used to evaluate them. (Please note that additional information is rapidly becoming available and the reader may want to update this information in the years following publication of this article). While the U.S. National Toxicology Program and the International Agency for Research on Cancer have classified the plasticizer DEHP as a possible human carcinogen, their methodologies have been criticized for potentially inaccurately ascribing results obtained with rodents to humans (141, 142). Mechanistic studies indicate that the carcinogenic response which DEHP produces in rodents is directly related to physiologic and metabolic changes that are specific to that species. Because the evidence indicates the response is an artifact to that species and not a true indication of human hazard, a number of regulatory bodies do not consider DEHP to pose a hazard to man. The Specialized Experts Working Group of the European Commission for instance, has concluded that there is no evidence to warrant the classification of DEHP as a carcinogen (143). DEHP is not regulated as a carcinogen by the U.S. Food & Drug Administration, which has long governed the plasticizer's use in medical devices and in food contact applications. DEHP-plasticized PVC is used in medical applications like blood bags where it is known to actually protect red blood cells from deterioration. Flexible PVC film is considered the most desirable material for wrapping meats, as it is oxygen-permeable and maintains the bright red color needed to make meats salable to consumers, as well as extending the shelf life of meats.

Fillers

Fillers are used to improve strength and stiffness, to lower cost, and to control gloss. The most common filler is calcium carbonate. It ranges in size from 0.07 to well over 50 μm . Some forms are treated with a stearic acid coating.

Clay fillers, such as calcined clay, improve electrical properties. Glass fibers, talc, and mica improve tensile strength and stiffness, but at a loss in ductility.

Pigments

A variety of pigments are added to PVC to give color, including titanium dioxide and carbon black.

Ultraviolet light stabilizers

One form of stabilization is to absorb the ultraviolet light. Both titanium dioxide and carbon black are strong ultraviolet light absorbers and effective in protecting the PVC. The carbon black is a stronger absorber than titanium dioxide and can therefore be used at lower levels in PVC for protection. For ultraviolet light absorption in transparent PVC or to improve pigmented systems, various derivatives of benzotriazole are used such as

2-[2'-hydroxy-3',5'-(di-t-butyl)phenyl]benzotriazole. Where tin carboxylate stabilizers are used instead of tin-mercaptide stabilizers, hindered amine light stabilizers, particularly with ultraviolet absorbers, are effective (144).

Biocides

Although PVC itself, and rigid PVC compounds are resistant to attack by microorganisms, plasticized PVC, in specific applications such as flashing and sealing boots on roofs, shower curtains, and swimming pools, may need protection. Many biocides, often containing arsenic compounds, are available for a balance of stability, compatibility, weatherability, and biocidal effectiveness.

Flame retardants

Since PVC contains nearly half its weight of chlorine, it is inherently flame retardant. Not only is chlorine not a fuel, but it acts chemically to inhibit the fast oxidation in the gas phase in a flame. When PVC is diluted with combustible materials, the compound combustibility is also increased. For example, plasticized PVC with >30% plasticizer, may require a flame retardant such as antimony oxide, a phosphate type plasticizer, or chlorinated or brominated hydrocarbons (145, 146).

Foaming or blowing agents

Cellular PVC can be made by a variety of techniques, such as whipping air into a plastisol, incorporating a gas under pressure or incorporating a physical blowing agent into the melt, or using a chemical blowing agent which releases a gas when it decomposes with heat.

The most common chemical blowing agent is 1,1'-azobisdicarbonyl, which decomposes with heat to release nitrogen gas. Typically the closed cell foams of rigid PVC range down to a density of 0.4 grams/cc. Physical blowing agents, such as chlorofluorocarbons which volatilize without changes in the chemical bonds, are capable of producing foams at down to 0.03 grams/cc. when used with a copolymer PVC. Because of the damage to the ozone layer in the stratosphere, not all chlorofluorocarbons are acceptable and newer types of physical blowing agents are becoming available which minimize ozone depletion.

PRODUCTION VOLUME AND ECONOMICS

Compound manufacture

All PVC must be compounded before it is used. For the wide variety of requirements where PVC performs, it is usually not feasible to maintain a technical compounding knowledge at the user level. Often processors and original equipment manufacturers (OEMs) will purchase compound, already balanced to meet the range of properties required and to do the best job economically. Some resin manufacturers are integrated to supply these compounds. Suppliers of merchant compound sold in North America are listed in figure 11 according to their approximate share (147). The total volume is not publicly available. These compounds are available in bulk railcars containing 80 tons (160,000 lbs.), in bulk trucks containing 20 tons (44,000 lbs.), boxes containing 600 kg (1200 - 1500 lbs.), or bags containing 23 kg (50 lbs.).

For businesses specializing in a large market with a limited product line, it can be economical to streamline the process, from compound manufacture to melt extrusion, to an extent that compounding can be integrated into the process. These businesses are not included in the following chart of merchant compound producers. Pipe and siding manufacturing are such processes where compounding is integrated in with the extrusion process. Other businesses that integrate compounding to some extent are window

manufacturing, barrier & packaging film manufacturing, and wire & cable
manufacturing.

Figure 11. North American merchant compound sales - 1995 (147).

int. = integrated PVC resin and compound manufacturing.

PVC resin manufacture

PVC resins are manufactured and sold for a wide variety of applications. About 14.5 billion pounds of resin are manufactured in North America and 53 billion pounds world wide. These resin manufacturers are broken down in figures 12 and 13. These resins are available in bulk railcars containing 80 tons (160,000 lbs.), in bulk trucks containing 20 tons (44,000 lbs.), boxes containing 600 kg (1200 - 1500 lbs.), or bags containing 23 kg (50 lbs.).

Figure 12. North American PVC resin capacity - 1995 - 6.6 billion Kgrams, (4.5 billion pounds) (1, 147, 148). Source for both fig 12 & 13: CMAI (148).

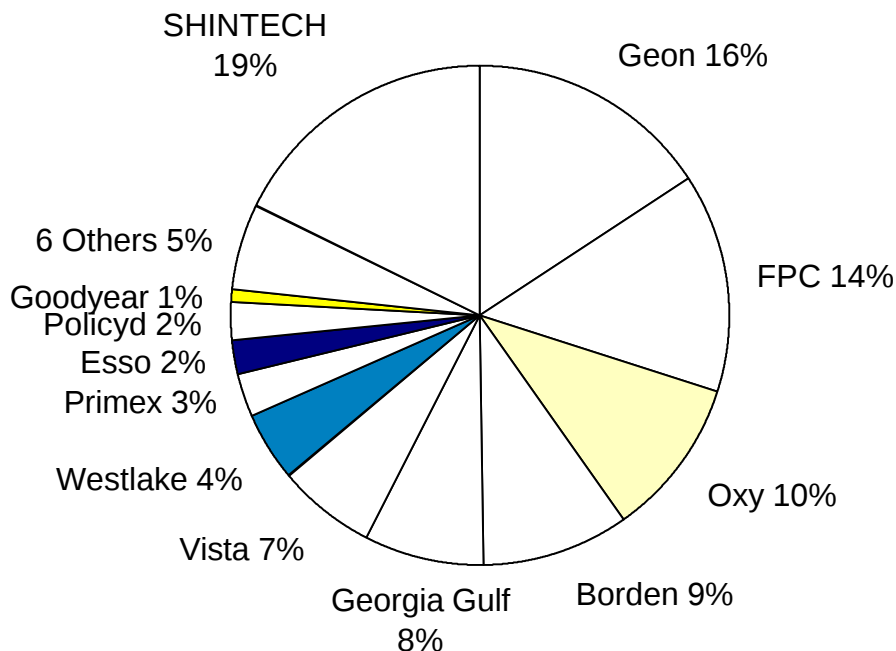
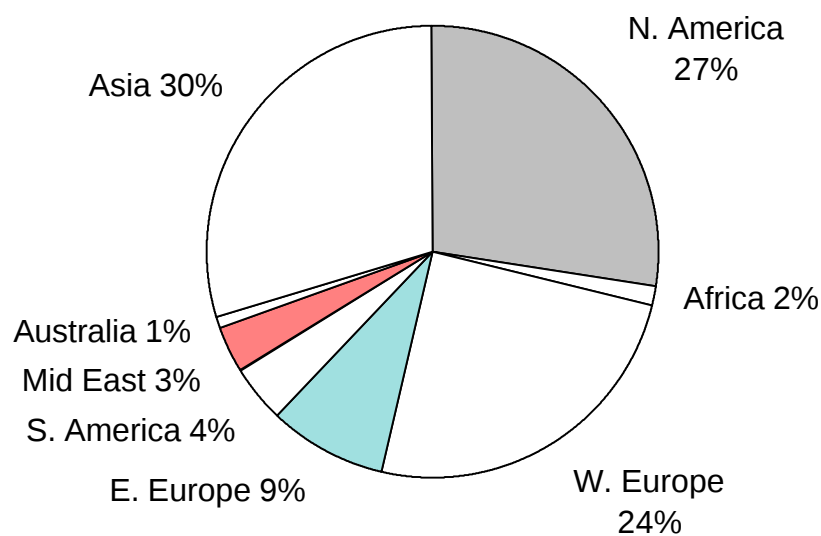


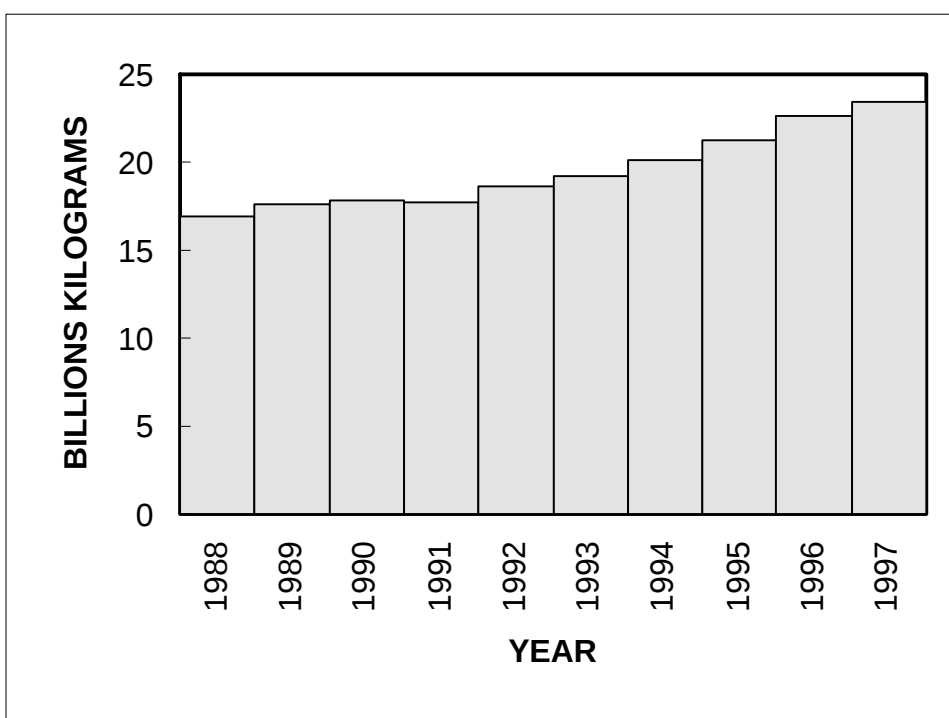
Figure 13. World PVC resin capacity - 1995 - 24 billion Kgrams (1, 147, 148).



World demand

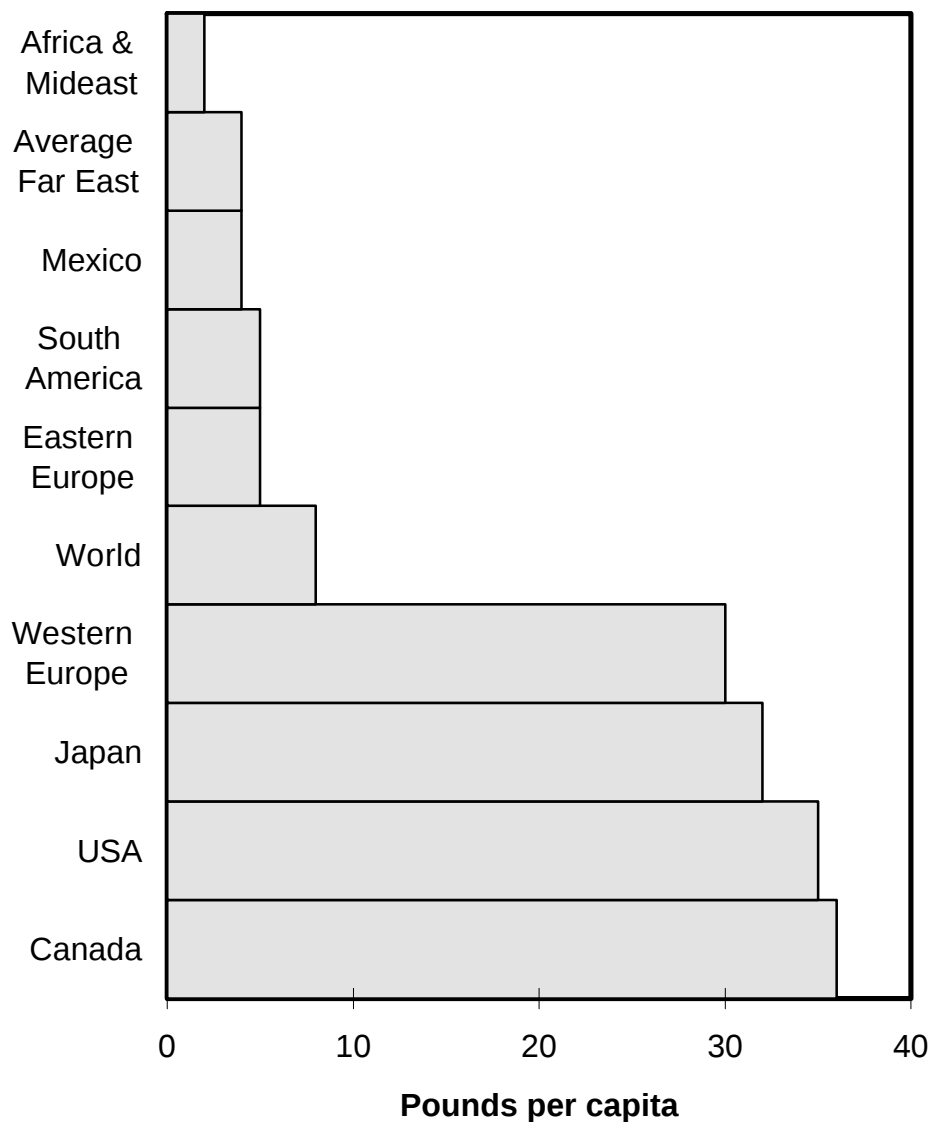
World demand is 22 billion kilograms (49 billion pounds) in 1995, with the growth accelerating (figure 14).

Figure 14. World demand for PVC. (1, 147, 148). Source: CMAI (148).



Per capita demand for PVC is high in Canada, USA, Japan, and Western Europe, with lots of room for growth in Africa, the Mideast, the Far East, Mexico, South America, and Eastern Europe (figure 15).

Figure 15. Per capita demand for PVC by region. (1, 147, 148). Source: CMAI
(148).



PVC resin price

PVC resin prices tend to be more stable than other plastics' prices, partly because only about half the molecule is based on hydrocarbon raw material

sources. Figure 16 compares PVC prices to other plastics' prices. Figure 17 looks at PVC's prices over several years.

Figure 16. PVC prices compared to other plastics' prices (147, 148). Source: CMAI (148).

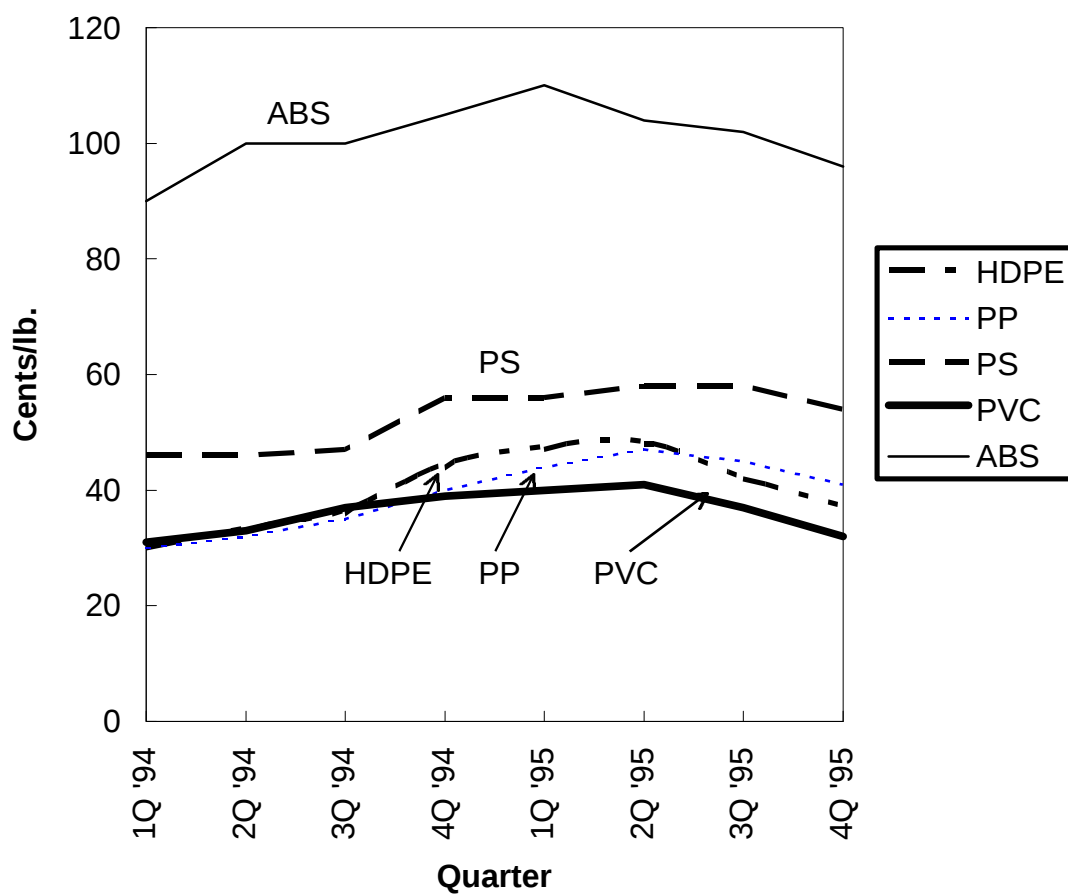
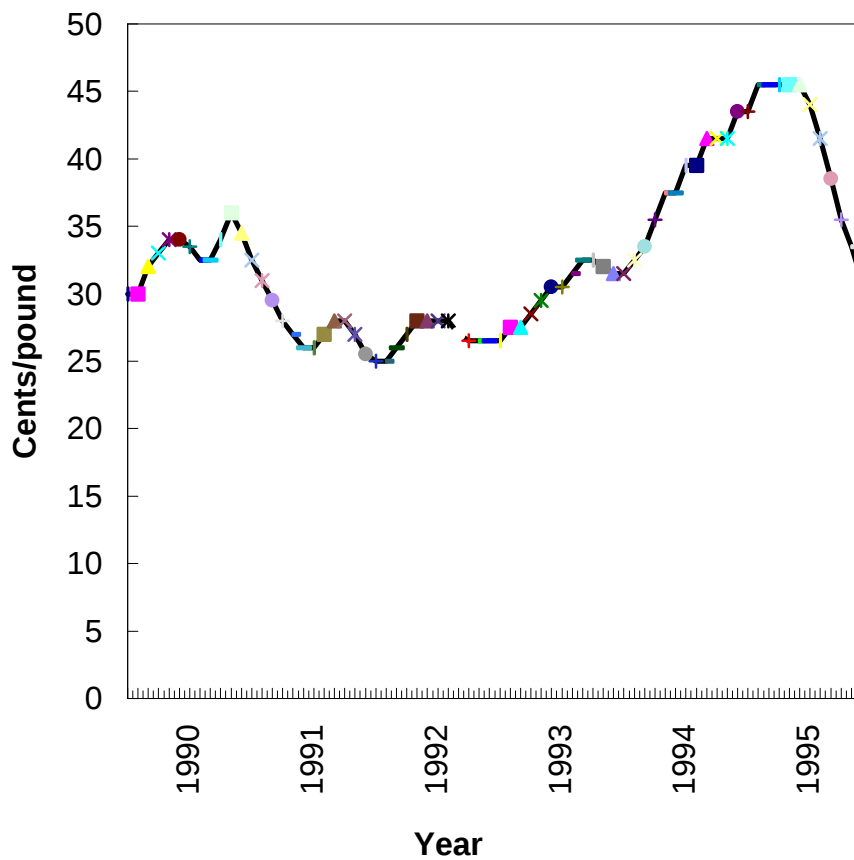


Figure 17. PVC prices over several years (147, 149, Chem Data, Inc.).



SPECIFICATIONS AND STANDARDS

Numerous specifications and standards are used to define PVC resins and compounds. Table 8 summarizes a few of these standards.

Table 8. Specifications and standards for PVC resins and compounds.

Standard	Description
ASTM D1755	PVC resin cell classification
ASTM D1784	PVC rigid compound cell classification
ASTM D1785	PVC pipe, schedule 40, 80, 120
ASTM D2287	PVC non-rigid (plasticized) compound cell classification
ASTM D2464	PVC THR fittings
ASTM D2466	PVC socket fittings, schedule 40
ASTM D2467	PVC socket fittings, schedule 80
ASTM D2665	DWV pipe and fittings
ASTM D2729	PVC Sewer pipe and fittings
ASTM D2740	PVC tubing
ASTM D2846	CPVC hot water distribution system
ASTM D2949	PVC DWV, 3 inch thin wall
ASTM D3033	PVC PSP sewer pipe and fittings
ASTM D3034	PVC PSM sewer pipe and fittings
ASTM D4216	PVC rigid building products compounds
NSF 14	PVC potable water, fittings
UL 62	PVC wire insulation
UL 83	PVC T, TW, THW, THWN, THHN wire insulation
UL 444	PVC telecommunications wire insulation jacket
UL 719	PVC NM jacket
UL 758	PVC AWM wire insulation
UL 1272	PVC TC cable jacket

USES AND MARKETS

PVC is so versatile that it can be compounded for a wide range of properties and it is used in a wide variety of markets. Most of the products are durable goods and have long lifetimes. Its use in short term, one time use, products is limited. A list of all major uses is found in table 9.

Table 9. USA Markets - 1995 (1).

		Metric tons
Construction		
	Flooring	120,000
	Pipe & Conduit	2,070,000
	Wire & Cable	180,000
	Siding	640,000
	Windows & Doors	140,000
	Extrusions	240,000
Packaging		
	Bottles	80,000
	Extrusions	160,000
	Calendering	390,000
Custom Moldings		50,000
Paste Processes		100,000
Textiles & Coating		70,000
Exports & Resale		850,000
Other		60,000
Total		5,360,000

Pipe and fittings

Pipe and fittings are a major market for PVC. These are a prime example of PVC as an engineering thermoplastic. These applications are designed with PVC for long term satisfactory performance. The applications are highly optimized for efficient production at high rates and minimal costs. Pipe manufacturing uses powder compounds, twin screw extruders, and vacuum sizing/cooling. Because of the volume of products, the process is highly developed. The products are designed to meet appropriate ASTM standards.

Products include pressure pipe of various sizes and drain, waste, and vent applications.

Chlorinated PVC (CPVC) is used in higher temperature applications such as for hot water piping. Because of its superior creep resistance, CPVC is also used in automated fire-safety sprinkler systems.

Weatherable siding, windows, & doors

PVC is accepted commercially as an excellent weathering material. Plastic materials are damaged by the sun, particularly by ultraviolet light. PVC's chemical response to weathering is well understood so that compounds and products can be designed for satisfactory performance outdoors. The mechanism of degradation starts by absorbing the sun's damaging ultraviolet light. This absorption is affected by PVC's previous thermal degradation during processing (150, 151). For this reason it is important to avoid thermal damage to PVC during processing. The absorbed light breaks bonds and forms free radicals. This leads to loss of hydrogen chloride and yellowing, and at the same time, oxidative bleaching (removal of yellowing). With a proper level of titanium dioxide to protect the PVC by its absorption of ultraviolet light (152), with a proper choice of the type of titanium dioxide to control the rate of the oxidation and bleaching process, and with a proper choice of other weather stable

ingredients, a well designed PVC formulation can be very durable to the weather.

As with pipe, siding manufacturing is highly optimized for efficient production at high rates and minimal costs. Siding uses powder compounds, twin screw extruders, and vacuum sizing/cooling. Because of the volume of products, the process is highly developed and optimized.

Other products such as windows, may or may not use the powder compound/twin screw extruder/vacuum sizing approach. For windows, this is a much more complex operation than for pipe, and requires large investments to develop and operate the process. Cubed compound, where the PVC grains are already broken down, can be run faster and on simpler single screw extruders, because lower melt temperatures are typical. Because elastic swell is reduced at lower melt temperatures, die design is simpler. Either vacuum sizing or air sizing/cooling is possible. The products are designed to meet appropriate ASTM standards. Products include siding, soffits, gutters & down spouts, windows, including the all-vinyl windows, and vinyl protected wood windows, and door glazing applications and garage doors.

Profiles

Complex profiles require specialty manufacturing skills to build, maintain, and operate extrusion dies, and cooling and sizing equipment that delivers the exact dimensions required. Cubed compound, where the PVC grains are already broken down, can be run faster and on simple single screw extruders, because low melt temperatures are typical. Because elastic swell is reduced at lower melt temperatures, die design is simpler and often is the same dimensions as the desired profile. Either vacuum sizing or air sizing/cooling is possible.

The low elastic swell is unique to PVC, and is due to the billion molecule flow units. Other plastics have higher die swell, which increases at lower melt temperatures (153). Thus those plastics have much more difficulty in designing dies and in maintaining a process to hold dimensions.

Wire & cable

PVC has been used in wire & cable applications since World War II, when the Navy demanded lower combustibility materials in construction. These products are manufactured by cross-head extrusion, usually from pellet compounds on single screw extruders. Some line speeds are 5000 feet/minute (60 mph). The compounds are optimized for the requirements, including low

temperature flexibility, high use temperature, especially low combustibility, weatherability, and high resistance to cut-through.

Injection molded products

Numerous housings, electrical enclosures, and cabinets are injection molded from rigid PVC. These take advantage of PVC's outstanding Underwriter Laboratories flammability ratings and easy molding into thin walled parts. PVC has developed melt flow capabilities to the point where it competes with essentially any other flame retarded engineering thermoplastic and molds easier than most.

Pipe fittings require quite high tensile and creep resistance, thus they are molded from compounds have less melt flow than for thin walled housings.

Plasticized compounds are injection molded into a variety of parts requiring elastomeric properties.

HEALTH AND SAFETY FACTORS (TOXICOLOGY)

MSDSs available

The potential health and safety effects of PVC and all PVC additives, as well as all feedstocks, are fully detailed by each manufacturer in material safety data sheets (MSDS), which are required by U.S. law. These are available to anyone who wishes to review them, and must be provided to all plant workers. While the processing and use of PVC is not any more hazardous than that of other plastic materials, this section goes into potential concerns in some detail.

Health [references (154 - 158)]

There are no significant health hazards arising from exposure to polyvinyl chloride at ambient temperature. However, a British study found a small decrease in breathing capacity for workers who smoked and were exposed to vinyl resin dust. This decrease was about one-seventh of that caused by normal aging and about equal to that expected with a one-pack-a-day cigarette smoker.

Since routine inhalation of dust of any kind should be avoided, reduction of exposure to polyvinyl chloride dust may be accomplished through the utilization of care when dumping bags, sweeping, mixing or doing other tasks

which can create dust. The use of an approved dust respirator is recommended where adequate ventilation may be unavailable.

At processing temperatures, most polymers emit fumes and vapors that may be irritating to the respiratory tract. This is also true for PVC and its additives, and this irritation may extend to the skin and eyes of sensitive people. Processing emissions exposure can also be greatly reduced or eliminated by the use of properly designed and maintained exhaust ventilation.

Decomposition of plastics, e.g., through greatly elevated temperatures above normal operating temperatures, can result in personnel exposure to decomposition or combustion products. In the case of PVC compounds, such decomposition would involve hydrogen chloride. Hydrogen chloride will cause irritation of the respiratory tract, eyes and skin. Depending upon the severity of exposure, physiological response will be coughing, pain and inflammation of the respiratory tract.

Fortunately, the pungent odor of hydrogen chloride provides an excellent warning signal, causing exposed personnel to be driven from the area which prohibits long term exposure. (The odor of hydrogen chloride is detectable as low as 1 - 5 ppm.)

First aid for exposure to processing emissions or decomposition products is simply to remove the exposed individual from the area of exposure. It is recommended that a physician examine those persons complaining of persisting irritation.

Fire and explosion

Polyvinyl chloride resin has a flash point of approximately 391°C (735°F) and a self-ignition temperature of approximately 454°C (850°F) - ASTM D-1929. In general, polyvinyl chloride burns with difficulty because a substantial amount of energy is required to break down the polymer into smaller fragments that will sustain combustion in the gas phase, principally as a consequence of the action of the halogen content of the material. Consequently, polyvinyl chloride is difficult to ignite. Fires will tend to extinguish naturally in the absence of a substantial external source of heat or flame. As mentioned previously, hydrogen chloride is generated during combustion. This action serves as a flame quencher in the vapor phase. Polyvinyl chloride will release less heat than many other combustible materials. Precautions should be taken similar to those for any other combustible materials, e.g., wood or other plastics.

Polyvinyl chloride powder has a very low tendency to explode. The minimum ignition energy for explosion is much higher than that of natural materials such as corn starch and flour and also exceeds those of other plastic

materials. However, as with any powder materials, care should be taken in addressing ignition sources in working and handling areas, should dusting occur. Also insure walkways and floors are cleared of polyvinyl chloride dust to prevent slippery footing.

With respect to firefighting where polyvinyl chloride is involved, water, ABC dry chemical or protein-type air foams should be used as extinguishing media. Carbon dioxide may be ineffective on larger fires due to lack of cooling capacity which may result in reignition. Firefighters should utilize a self-contained breathing apparatus (SCBA) in positive-pressure mode. In addition, for an enclosed or poorly ventilated area, an SCBA should be worn during cleanup immediately after a fire as well as during the attack phase of firefighting operations.

PVC should not be melt mixed with acetal polymers. These polymers are chemically incompatible; mixing could cause rapid decomposition and gas evolution.

Toxicology

Toxicology studies have shown polyvinyl chloride to be equivocally tumorigenic through oral and implant studies. The International Agency for Research on Cancer (IARC) shows inadequate evidence that polyvinyl chloride is carcinogenic in animals or humans and has an overall evaluation of 3 (not

classifiable). The United States has no occupational exposure limits for polyvinyl chloride except as particulates not otherwise classified (PNOC). For PNOC's the American Conference of Governmental Industrial Hygienists (ACGIH) has a threshold limit value (TLV) of 10 mg/m³ for inhalable and 3 mg/m³ for respirable while the Occupational Safety and Health Act (OSHA) permissible exposure limits (PEL's) are 15 mg/m³ for total dust and 5 mg/m³ for the respirable fraction.

Regulatory

Polyvinyl chloride is listed on the TSCA inventory and the Canadian Domestic Substances List (DSL) as ethene, chloro-, homopolymer [CASRN: 9002-86-2]. Since polymers are not listed on the European Community Commercial Chemical Substances listing or EINECS, polyvinyl chloride is listed through its monomer, vinyl chloride [CASRN: 75-01-4]. In the United States polyvinyl chloride is an EPA Hazardous Air Pollutant under the Clean Air Act Section 112 (40 CFR 61) and is covered under the New Jersey Community Right-to-Know Survey: N.J. Environmental Hazardous Substances (EHS) List as CHLOROETHYLENE, POLYMER with a reporting threshold of 500 lbs.

ENVIRONMENTAL CONSIDERATIONS AND RECYCLING (ref. 159-164)

Chlorine, the material used to make PVC

Chlorine is the 20th most common element on Earth, found virtually everywhere: in rocks, oceans, plants, animals and human bodies. It is essential to human life. Free chlorine is produced geothermally within the earth, and occasionally finds its way to the earth's surface in its elemental state (165). More usually, however, it reacts with water vapor to form hydrochloric acid.

Hydrochloric acid reacts quickly with other elements and compounds, forming stable compounds (usually chlorides) such as sodium chloride (common salt), magnesium chloride and potassium chloride, all found in large quantities in sea water.

The chlorides found in common salt water are an essential element in all body fluids, and in this form, makes up about 0.15 percent of total body weight. A large number of complex organic chlorides and organochlorines are naturally produced chemicals, widely present in nature, and play many essential roles (166 - 169). Chlorine is also an essential element of naturally occurring anti-bacterial and anti-fungal agents such as chlortetracycline, chloramphenicol and griseofulvin, which have revolutionized the treatment of human bacterial and fungal infections.

Even more complex cyclic organochlorines (including dioxins and furans) are produced from burning wood and other vegetable matter, and are natural by-products of forest fires (170, 171). Chlorine-based chemicals are everywhere, and have been so since before the existence of man. Both dioxins and furans have been found in lake sediments dating back to 1860 (172) and samples taken from ice cores in Greenland dating back to 1869 show definite "spikes" in chlorine content correlating with volcanic activity (173). Given the fact that chlorine compounds are naturally produced in such vast quantities, and so widely distributed in the natural world, banning production of chlorine to keep chlorine compounds out of the environment would be futile, extremely costly, and would deprive the world of hundreds of products critical to society's health and well-being. And the environment would be worse off.

Over 30% of the chlorine produced on a global basis goes to make PVC. Not only is chlorine essential to the chemical composition of PVC, it provides a number of unique properties that give this versatile plastic a distinct advantage in product applications and the marketplace. It makes PVC inherently flame retardant. PVC is the world's leading electrical material, with over 500 million pounds used annually for wire and cable insulation and sheathing, electrical conduit, boxes and components. PVC is over 50% chlorine and as a result, is one of the most energy efficient polymers. Chlorine makes PVC far more environmentally acceptable than other materials that are totally dependent on petrochemical feedstocks. In addition, recycling PVC is easier because the

chlorine in PVC acts as a marker, enabling automated equipment to sort PVC containers from other plastics in the waste stream (174).

Vinyl Solid Waste and Recycling

Although vinyl is the world's second most widely used plastic, less than one-half percent by weight is found in the municipal solid waste stream. Most of that consists of vinyl packaging, bottles, blister packaging and flexible film. That's because most vinyl applications are long-term uses, such as pipe and house siding, that are not disposed of quickly. Vinyl wastes are handled by all conventional disposal methods: recycling, landfilling and incineration (including waste-to-energy).

Vinyl is recycled by at least 170 recyclers in the U.S. and Canada. In 1994, about 6.5 million pounds of post-consumer vinyl were recycled in the U.S. An estimated additional 300 million pounds of vinyl post industrial scrap was diverted from landfills and recycled into second-generation products. More than 3,500 communities accept vinyl products in their recycling programs, which is about 25 percent of all communities that recycle.

In landfills, vinyl wastes, like all plastics, are extremely resistant to decomposition. In fact, high-technology landfills are often lined with thick-gauge

vinyl and use PVC pipe to handle liquid leachate and methane gas, to protect the environment.

Vinyl compares favorably to other packaging materials. In 1992, a lifecycle assessment comparison of specific packages made from glass, paperboard, paper and selected plastics concluded that vinyl was the material with the lowest production energy and carbon dioxide emissions, and the lowest fossil fuel and raw material requirements of the plastics studied (175). Vinyl saves more than 34 million BTUs per 1,000 pounds manufactured compared to the highest energy-consuming plastic (176).

Incinerating PVC Wastes

A recent study sponsored by the American Society of Mechanical Engineers (ASME), involving the analysis of over 1,700 test results from 155 large-scale, commercial incinerator facilities throughout the world, found no relationship between the chlorine content of waste and dioxin emissions from combustion processes. Instead, the study stated, the scientific literature is clear that the operating conditions of combustors are the critical factor in dioxin generation (177). This work includes and confirms a number of other studies, most notably, the work conducted in 1987 by the New York Energy Research and Development Authority. Those tests revealed that the presence or absence

of PVC had no effect on the amount of dioxin produced during the incineration process (178).

Incinerator scrubbing systems can remove about 99 percent of the hydrogen chloride generated by incinerating vinyl plastics and other chlorine-containing compounds and materials (179). New requirements from the U.S. Environmental Protection Agency make scrubbers mandatory on all incinerators so that they can neutralize a range of acid gases, including sulfur dioxide and nitrogen oxide, which are produced by a variety of materials. Since acid generated in incinerators comes from a variety of sources including such things as table salt and paper products, scrubbers are necessary whether or not PVC is present in the waste feed (180). A recent study, conducted by Midwest Research Institute and published by the ASME, concluded that removing vinyl from the waste stream would not eliminate the need either for air pollution control devices and monitoring equipment, nor would it influence the choice of incineration equipment (181).

Municipal incinerators are often targeted as a primary cause of acid rain. In fact, power plants burning fossil fuels, which produce sulfur dioxide and nitrogen oxide, are actually the leading cause of acid rain, along with automotive exhaust (182, 183). In Europe and Japan, studies show that only about 0.02 percent of all acid rain can be traced to incineration of PVC (184).

Common misperceptions about vinyl

a) No one is recycling vinyl.

Not true. Of course, industrial scrap vinyl has been recycled for years, but now, post-consumer vinyl recycling is growing, too, with about 6.5 million pounds of post-consumer vinyl (primarily bottles) currently being recycled. When the Council for Solid Waste Solutions (now, the American Plastics Council) conducted a nationwide survey in 1991, it found that there were an estimated 1,100 municipal recycling programs in place or planned in the United States that include vinyl.

b) There's no market for recycled vinyl.

Not true. In 1989, the University of Toledo identified nearly 100 uses for recycled vinyl. Overall, the potential demand for recycled vinyl is estimated to be over twice the potential supply of all vinyl bottles produced in the US each year (494 million pounds needed vs. 207 million pounds available via recycling of bottles). A recent directory published by the Vinyl Institute lists nearly 50 companies that make commercial products out of recycled vinyl (185).

c) Vinyl is the problem in municipal recycling because it contaminates other resins.

Not true. Contamination occurs whether or not vinyl is present. Other resins are just as much a contamination problem for vinyl. Except for commingled plastics applications, different plastic materials cannot be mixed successfully in most recycled products applications. This is why it's so important to efficiently separate one plastic from another. Thanks to the chlorine that is present in it, vinyl lends itself very well to automated sorting technology.

d) PET and HDPE packaging are listed as 1 and 2 in the SPI recycling coding system because they are the most recyclable.

Not true. The numbers assigned to each plastic in the SPI coding system are purely arbitrary and do not reflect the material's recyclability.

e) Vinyl gives off dioxin when it's incinerated.

Misleading. A study conducted by ASME in 1995, found that the presence--or absence--of chlorine containing wastes in incinerators had no effect on the levels of dioxin produced. Rather, it was found that incinerator operating conditions (primarily temperature) were the key to controlling dioxin formation (168). Most recently, German officials examined the issue of incinerating vinyl waste and decided there was no cause for concern.

f) Vinyl should be banned from incinerators because it contains heavy metal additives.

Misleading. This is an evolving issue. Most vinyl products do not contain heavy metals and vinyl is a small fraction in feed to incinerators. Re-formulation to replace heavy metals is in progress but some use will continue. Banning vinyl from incinerators will not eliminate this problem. Rather, regulations should specify that incinerator residues (ash) be disposed of appropriately.

g) European packagers, grocery stores, and regulators have banned vinyl.

Not true. There is only one ban on vinyl packaging in Europe, and that is on the use of vinyl bottled mineral water in Switzerland, a commercial ploy to block the sale of French mineral water in that country. There is a voluntary agreement in Denmark by industry to substitute alternatives to vinyl packaging when feasible. Some municipalities in Germany have restricted the use of certain vinyl products in municipally funded building projects. Industry is working to change those restrictions with several notable reversals, including Berlin and Bielefeld. Government studies on PVC in Belgium and The Netherlands concluded there should be no bias against the use of PVC (186 - 188). Elsewhere in Europe, vinyl packaging continues to be widely used. In Great Britain, one of the leading retailers, Marks & Spencer, has chosen vinyl over other materials as the chain's most "environmentally friendly polymer." In Switzerland, retailer Migros has stated that its whole attitude toward vinyl will change when incinerator scrubber technology is fully employed. The current trend in Europe (led by the Germans) is to take a comprehensive look at waste

reduction and make industry a partner in that process. This involves all industries, not just the vinyl industry. Overall, Europe remains a larger consumer of vinyl packaging than the U.S.

h) Vinyl plastics decompose in landfills and give off vinyl chloride monomer.

Not true. Like all plastics, vinyl is an extremely stable landfill material. It resists chemical attack and degradation, and is so resistant to the conditions present in landfills it is often used to make landfill liners. On those occasions when vinyl chloride monomer is detected in landfills, it typically can be traced to the presence of other chemicals and solvents.

i) Other plastics are more environmentally "friendly" than vinyl.

Not true. A recent study conducted by Chem Systems, Inc., an independent consulting firm, compared vinyl to a number of other packaging materials and found that vinyl consumed the least amount of energy, used the lowest level of fossil fuels, consumed the least amount of raw materials and produced the lowest levels of carbon dioxide of any of the plastics studied. In fact, the Norwegian environmental group Bellona has concluded that "a generally reduced use of vinyl plastics will, given today's circumstances, lead to a worsening of the environmental situation."

j) In a fire, vinyl is unusually hazardous and damaging

Not true. The real hazards in a fire are carbon monoxide and heat; these are especially a problem with other materials that readily burn. Because vinyl products contain chlorine, they are inherently flame-retardant and resist ignition. When it does burn however, vinyl produces carbon monoxide, carbon dioxide

and hydrogen chloride. Of these, the most hazardous is carbon monoxide.

Hydrogen chloride is an irritant gas that can be lethal at extremely high levels.

However, research indicates that those levels are never reached or even approached in real fires. All organic materials, when burned, release a lengthy list of chemical by-products. For instance, when wood burns, as many as 175 different fire gases may be produced, including benzene and acrolein (189).

Burning wool will produce hydrogen cyanide. Even the simple act of barbecuing a steak or smoking a cigarette will produce dioxin (190). More importantly, virtually all burning materials produce carbon monoxide, which is by far considered the greatest toxic hazard in fires because of the abundant levels produced and the low levels that cause death (191).

The U.S. fire death rate is decreasing, dropping from a rate of 76 per million in the 1940s (when most construction and decorative products were made of "natural" materials), to 29 per million in the 1980s (by which time, PVC had replaced natural materials in numerous applications) (192). This downward trend can be attributed in large part to improved building codes and the broader use of sprinkler systems and smoke detectors. However, the increased use of more fire-resistant materials, like PVC, deserves part of the credit for this improvement.

Hydrogen chloride is produced when PVC burns. A series of tests for the Federal Aviation Administration studied this issue. In those studies, test animals were able to survive exposures to hydrogen chloride reaching 10,000 ppm(193). More recent studies indicate less of a potential for delayed effects on lung

function than expected (194). In a typical fire, hydrogen chloride levels rarely exceed 300 ppm, a fact confirmed by the Boston Fire Department and Harvard University (195). In hundreds of autopsies conducted on fire victims in the U.S., not one death has been linked to the presence of PVC.

Bell Laboratories studied wire and cable compounds made of PVC or other halogen-based compounds vs. halogen-free compounds and found that neither type of material presented a clear-cut advantage in a fire, and that the halogenated compounds sometimes outperformed the nonhalogenated products in terms of creating less corrosion (196).

ACKNOWLEDGMENT

I wish to thank The Geon Company for allowing me over the years to build a knowledge of PVC that makes this review possible. I wish to thank Lu Liu and Dick Roman for assistance in determining the size of this industry, Thanks to Mark Hross for his assistance on the section on health and safety, and thanks to D'Lane Wisner and Fred Krause for helping with the environmental considerations and recycling. Of course the previous authors of this review, John Davidson and the late Keith Gardner, made my job easier.

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Search date: November, 1995.

ABSTRACT

PVC has a large sales volume, second only to polyethylene. Its high chlorine content provides it with a very high level of combustion resistance for building products, electrical enclosures, and wire & cable insulation. PVC has a unique ability to be compounded with a wide variety of additives, making it possible to produce materials in a range from flexible elastomers (the first thermoplastic elastomers) to rigid compounds, materials that are weatherable such as for siding and windows, compounds that have stiff melts and little elastic recovery for outstanding dimensional control useful in profile extrusion, or low viscosity melts which compete effectively with ABS and PC/ABS in thin walled injection molding parts such as computer monitor housings.

Some of PVC's properties are attributed to unique morphology. The polymer precipitates from its monomer and grows into primary particles, which are later the melt flow units. Fusion into larger structures and product strength are controlled by break-down of the grains into primary particles, by the choice of additives, by the amount of melting (temperature), and by the number of tie molecules (molecular weight).

The main type of polymerization is the suspension process, with significant polymerization made by the mass process, solution process, micro-suspension, and emulsion process. In the suspension process, the polymerization takes place in droplets of monomer suspended in water.

From an environmental viewpoint, PVC is over 50% chlorine and as a result, is one of the most energy efficient polymers, makes PVC inherently flame retardant, and acts as a marker, enabling automated equipment to sort PVC containers from other plastics in the waste stream. Vinyl is recycled by at least 170 recyclers in the U.S. and Canada and more than 3,500 communities accept vinyl products in their recycling programs. The analysis from 155 large-scale, commercial incinerator facilities, found no relationship between the chlorine content of waste nor the addition of PVC, and dioxin emissions from combustion processes. New requirements from the U.S. EPA make scrubbers mandatory on all incinerators and are necessary whether or not PVC is present in the waste feed.